Comparison of perioperative costs with fast-track vs standard endovascular aneurysm repair

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Background: Perioperative health care utilization and costs in patients undergoing elective fast-track vs standard endovascular aneurysm repair (EVAR) remain unclear.

Methods: The fast-track EVAR group included patients treated with a 14 Fr stent graft, bilateral percutaneous access, no general anesthesia or intensive care monitoring, and next-day hospital discharge. The standard EVAR group was identified from Medicare administrative claims using a matching algorithm to adjust for imbalances in patient characteristics. Hospital outcomes including operating room time, intensive care monitoring, hospital stay, secondary interventions, and major adverse events (MAEs). Perioperative outcomes occurring from hospital discharge to 30 days postdischarge included MAE, secondary interventions, and unrelated readmissions.

Results: Among 1000 matched patients (250 fast-track; 750 standard), hospital outcomes favored the fast-track EVAR group, including shorter operating room time (2.30 vs 2.83 hrs, \(P<0.001\)), shorter hospital stay (1.16 vs 1.69 d, \(P<0.001\)), less need for intensive care monitoring (4.4% vs 48.0%, \(P<0.001\)), and lower secondary intervention rate (0% vs 2.4%, \(P=0.01\)). Postdischarge outcomes also favored fast-track EVAR with a lower rate of MAE (0% vs 7.2%, \(P<0.001\)) and all-cause readmission (1.6% vs 6.8%, \(P=0.001\)). The total cost to the health care system during the perioperative period was $26,730 with fast-track EVAR vs $30,730 with standard EVAR. Total perioperative health care costs were $4000 (95% CI: $3130–$4830) lower with fast-track EVAR vs standard EVAR, with $2980 in savings to hospitals and $1030 savings to health care payers.

Conclusion: A fast-track EVAR protocol using a 14 Fr stent graft resulted in shorter procedure time, lower intensive care utilization, faster discharge, lower incidence of MAE, lower readmission rates, and lower perioperative costs compared to standard EVAR.

Keywords: abdominal aortic aneurysm, cost, EVAR, fast-track, Medicare, percutaneous, perioperative

Introduction

Endovascular aneurysm repair (EVAR) has become the first-line treatment for abdominal aortic aneurysm (AAA) requiring intervention at most centers. Compared to open surgical AAA repair, EVAR is associated with lower risk for early mortality and morbidity.\(^1\) However, EVAR remains costlier than surgical repair\(^2\) and results in negative operating margins for hospitals.\(^3\) In 2015, the Centers for Medicare and Medicaid Services (CMS) increased EVAR reimbursements by 14% for uncomplicated cases and 24% for cases with major comorbidity or complication,\(^4\) but these increases are insufficient to make EVAR a profitable procedure at most hospitals. In the current economic environment with bundled payments\(^5\) and financial
penalties for higher than average readmission rates, adop-
tion of treatment pathways that improve operational ef-
ciency without compromising patient safety may potentially
improve hospital profit margins.

Enhanced recovery after surgery pathways, also known
as fast-track surgery, have been implemented across many
medical disciplines, including vascular surgery. Over
the last decade, several studies have evaluated the feasibility
of fast-track EVAR in well-selected patients. While the
elements of these programs differ slightly, the general
premise involves regional/local anesthesia, percutaneous
vascular access, no intensive care monitoring, and/or
early discharge. While cost reductions with fast-track
EVAR have been reported, these studies were performed
outside the US or evaluated outpatient EVAR, a pro-
cedure not currently eligible for reimbursement in the US.
Thus, the applicability of these findings to the US health
care system is unclear. The purpose of this study was to
compare perioperative health care utilization and costs
from a hospital and health care payer perspective in
patients undergoing fast-track EVAR vs matched patients
undergoing standard EVAR.

**Methods**

**Data sources**

The fast-track EVAR population included patients enrolled
in the prospective multicenter LIFE registry of elective
EVAR utilizing bilateral percutaneous access with a 14
Fr stent graft (Ovation Prime, Endologix, Inc., Irvine,
CA), no general anesthesia and intensive care monitoring,
and next-day hospital discharge. A total of 250 patients
were enrolled at 31 centers in the United States from
October 2014 to May 2016. The study was prospectively
registered at ClinicalTrials.gov (NCT02224794). The
study design and perioperative outcomes from this
study have been presented elsewhere. Patients were
enrolled in consecutive fashion and were eligible for the
study if they had a AAA anatomically suitable for elective
EVAR and were deemed by the treating physician to be
appropriate for a fast-track EVAR protocol based on pre-
defined study eligibility criteria (Table 1).

After obtaining approval from Western Institutional
Review Board (Puyallup, WA) with a waiver of consent
given the use of deidentified claims data, a standard EVAR
population was identified using administrative claims on all
Original Medicare beneficiaries available from CMS.
Medicare is the primary health care payer in 79% of
EVAR procedures; thus, the data derived from Medicare
claims are representative of nationwide EVAR outcomes.

We included all hospital claims submitted to the Inpatient
Prosp ective Payment System and Outpatient Prospective
Payment System, which were linked by beneficiary in
blinded fashion across the continuum of care.

The standard EVAR population included adult Medicare
beneficiaries who underwent elective EVAR between January
2014 and August 2015. The latter date was selected to allow
for follow-up through September 2015, which corresponds to
the date of transition from International Classification of
Diseases, Ninth Revision, Clinical Modification (ICD-9-CM)
to ICD-10-CM codes. Patients were identified using a combi-
nation of Medicare Severity-Diagnosis Related Group (MS-
DRG) codes and ICD-9-CM codes. We initially included
patients with MS-DRG code 237 (Major cardiovascular
procedures with major complication or comorbidity or thoracic
aortic aneurysm repair) or 238 (Major cardiovascular
procedures without major complication or comorbidity), ICD-9-CM
diagnosis code of 441.4 (Abdominal aneurysm without men-
tion of rupture), and ICD-9-CM procedure code of 39.71
(Endovascular implantation of other graft in abdominal
aorta). We then excluded patients with any of the following
ICD-9-CM diagnosis codes: 441.0 (Dissection of aorta), 441.1
(Thoracic aneurysm, ruptured), 441.2 (Thoracic aneurysm
without mention of rupture), 441.3 (Abdominal aneurysm
ruptured), 441.5 (Aortic aneurysm of unspecified site, rup-
tured), 441.6 (Thoracoabdominal aneurysm ruptured), 441.7
(Thoracoabdominal aneurysm without mention of rupture),
441.9 (Aortic aneurysm of unspecified site without mention
of rupture), 421.0 (Acute and subacute bacterial endocarditis),
or 759.82 (Marfan syndrome). In order to account for the main
exclusion criteria that were applied in the fast-track EVAR
study, we also excluded patients with congestive heart failure,
renal insufficiency, unstable angina, critical limb ischemia,
major surgery, or reintervention procedure 30 days prior to
the EVAR procedure, myocardial infarction, or stroke within
the last 3 months, and patients with life expectancy less than 1
year estimated by a Charlson comorbidity index of 5 or
greater. Exclusion conditions in the standard EVAR group
were identified from Medicare claims in the 12-month period
prior to the procedure.

**Patient matching**

In order to adjust for remaining imbalances in baseline patient
characteristics between fast-track vs standard EVAR groups,
we applied a 3-to-1 (standard-to-fast-track) algorithm that
matched patients on key characteristics, including age, sex,
chronic obstructive pulmonary disease, diabetes mellitus, coronary artery disease, valvular heart disease, cardiomyopathy, myocardial infarction, hypertension, peripheral vascular disease, and carotid artery disease. Matching was performed using the nearest neighbor algorithm that optimized the number of characteristics shared between groups.

Health care utilization
Hospital outcomes included operating room time, need for intensive care monitoring, length of hospital stay, secondary intervention, and major adverse event (MAE). An MAE was defined as myocardial infarction, stroke, renal failure, respiratory failure, paralysis, or bowel ischemia. Secondary interventions included endovascular or open surgical reinterventions using the coding scheme of Edwards and colleagues. Perioperative outcomes occurring over 30 days postdischarge included MAE, secondary interventions, and readmissions for reasons other than secondary intervention.

Health care cost estimation
Health care utilization costs were obtained from Medicare Hospital Standard Analytic Files. Costs were estimated

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<table>
<thead>
<tr>
<th>Table 1 Main study entry criteria for patients treated with fast-track endovascular aneurysm repair</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Main inclusion criteria</strong></td>
</tr>
<tr>
<td>• Age ≥18 years</td>
</tr>
<tr>
<td>• Male or non-pregnant female</td>
</tr>
<tr>
<td>• Candidate for elective open surgical AAA repair</td>
</tr>
<tr>
<td>• AAA &gt;5.0 cm diameter, increased ≥0.5 cm diameter in last 6 months, or maximum diameter exceeding 1.5 times its normal diameter</td>
</tr>
<tr>
<td>• Suitable anatomy for endovascular repair with the Ovation Prime Stent Graft</td>
</tr>
<tr>
<td>• Suitable anatomy to allow Perclose ProGlide Suture-Mediated Closure System via the pre-close technique, including:</td>
</tr>
<tr>
<td>o ≥5 mm in diameter,</td>
</tr>
<tr>
<td>o At least 2 cm segment for access, 10 mm above the origin of the profunda femoris branch, and 10 mm below the lower margin of the inferior epigastric artery as determined on preoperative contrast-enhanced CT, angiography, or ultrasound,</td>
</tr>
<tr>
<td>o No calcification on the anterior wall or circumferential (&gt;50%) calcification on the posterior wall,</td>
</tr>
<tr>
<td>o No prior groin incision, hematoma, or significant scarring,</td>
</tr>
<tr>
<td>o No prior clip or collagen-based vascular closure device placement within 90 days of the procedure,</td>
</tr>
<tr>
<td>o No prior femoral artery needle puncture within 30 days of the procedure, and</td>
</tr>
<tr>
<td>o No current active localized groin infection, traumatic vascular injury, femoral artery aneurysm, arteriovenous (AV) fistula, or pseudoaneurysm</td>
</tr>
<tr>
<td><strong>Main exclusion criteria</strong></td>
</tr>
<tr>
<td>• Dissecting or acutely ruptured AAA</td>
</tr>
<tr>
<td>• Acute vascular injury</td>
</tr>
<tr>
<td>• Prior AAA or iliac artery repair</td>
</tr>
<tr>
<td>• Mycotic AAA or active systemic infection</td>
</tr>
<tr>
<td>• Unstable angina</td>
</tr>
<tr>
<td>• Unstable peripheral artery disease with critical limb ischemia</td>
</tr>
<tr>
<td>• Congestive heart failure (NYHA class III or IV)</td>
</tr>
<tr>
<td>• Myocardial infarction or stroke within the past 3 months</td>
</tr>
<tr>
<td>• Need for renal artery coverage (e.g., Chimney graft)</td>
</tr>
<tr>
<td>• Planned adjunctive devices (e.g., renal stent)</td>
</tr>
<tr>
<td>• Major surgery or interventional procedure within the past 30 days</td>
</tr>
<tr>
<td>• Connective tissue disease (e.g., Marfan’s or Ehler’s–Danlos syndrome)</td>
</tr>
<tr>
<td>• History of bleeding disorder or refuses blood transfusions</td>
</tr>
<tr>
<td>• Dialysis-dependent renal failure or serum creatinine &gt;2.0 mg/dL</td>
</tr>
<tr>
<td>• Morbid obesity (BMI ≥40 kg/m²)</td>
</tr>
<tr>
<td>• Home oxygen use</td>
</tr>
<tr>
<td>• Patient admitted from skilled nursing facility</td>
</tr>
<tr>
<td>• Life expectancy &lt;1 year</td>
</tr>
<tr>
<td>• Anticipated inability to discharge patient within 1 day</td>
</tr>
<tr>
<td>• Participation in investigational device or drug clinical trial</td>
</tr>
<tr>
<td>• Intolerance/hypersensitivity to anticoagulation, contrast media, or stent graft components</td>
</tr>
</tbody>
</table>
from a hospital perspective, a health care payer perspective, and a total health care utilization perspective. The hospital perspective included hospital utilization and costs. Hospital costs were estimated from hospital charges multiplied by the hospital’s cost-to-charge ratio. The health care payer perspective included perioperative utilization occurring over 30 days postdischarge, with costs adjusted for health care payer mix. Costs were adjusted to represent national averages by assuming a nationwide primary health care payer mix of 83% Medicare and 17% private payers. This was a reasonable approximation since Medicare and private payers account for 95% of the primary health care payers in elective EVAR cases.15 We estimated private payer costs by applying a private payer to Medicare payment ratio for cardiovascular procedures.19 The total health care utilization perspective was calculated as the sum of hospital costs and health care payer costs.

**Data analysis**

The cost of individual elements contributing to overall hospital cost was estimated by multivariable linear regression. Total hospital costs were regressed on hospital length of stay and indicator variables for intensive care monitoring, MAE, and secondary intervention. Residuals plots were assessed for evidence of non-linearity or bias. Costs were not adjusted for inflation since all cases in both groups were performed over a 20-month period. Probability parameters were modeled using a beta distribution and continuous parameters were modeled with a normal distribution. The private payer to Medicare payment ratio was modeled with a triangular distribution. For outcomes with no observed events, we used a beta distribution with mean of 0.1% to accommodate model requirements. Monte Carlo probabilistic sensitivity analysis with 10,000 iterations examined the effect of combined uncertainty across all model parameters. Additional sensitivity analyses were performed to assess the robustness of the base-case results. Cost models were developed using TreeAge Pro 2017 (TreeAge Software, Williamstown, MA).

**Results**

**Patient characteristics**

Before matching, we identified 10,429 patients electively treated with standard EVAR. Compared to the fast-track EVAR group, these patients were older with more comorbidities. Following matching, this study included 1000 patients treated with elective EVAR—250 with fast-track EVAR and 750 with standard EVAR. Baseline patient characteristics were well matched between groups, with exceptions of higher incidence of renal insufficiency (10% vs 0%) and congestive heart failure (4% vs 0%) in the fast-track EVAR group. Since creatinine >2.0 mg/dL, or New York Heart Association class III or IV were exclusion criteria in the fast-track EVAR study, patients with renal insufficiency and creatinine ≤2.0 mg/dL, or CHF classified as NYHA class I or II remained eligible. Therefore, we conservatively excluded patients with renal insufficiency and congestive heart failure from the matched standard EVAR group (Table 2). Mean aortic characteristics in the fast-track EVAR group included AAA diameter of 51±8 mm, proximal neck angle of 24±17 degrees, proximal neck length of 24±14 mm, and external iliac diameter of 7.4±1.7 mm. Aortic characteristics were not available in Medicare claims for the standard EVAR group.

**Perioperative clinical outcomes**

Hospital outcomes favored the fast-track EVAR group, including shorter operating room time (2.30 vs 2.83 hr, \(P<0.001\)), shorter hospital stay (1.16 vs 1.69 d, \(P<0.001\)), less need for intensive care monitoring (4.4% vs 48.0%, \(P<0.001\)), and lower secondary intervention rate (0% vs 2.4%, \(P=0.01\)). Postdischarge outcomes generally favored fast-track EVAR with a lower rate of MAE (0% vs 7.2%, \(P<0.001\)) and all-cause readmission (1.6% vs 6.8%, \(P=0.001\)). Utilization and cost estimates for all model parameters are reported in Table 3. The all-cause readmission rate in the fast-track EVAR group was 1.6% (4/250); readmissions were attributable to renal insufficiency (0.4% [1/250]), cardiomyopathy (0.4% [1/250]), diverticulosis (0.4% [1/250]), and chronic obstructive pulmonary disease (0.4% [1/250]). In matched patients undergoing standard EVAR, the all-cause readmission rate was 6.8% (51/750). The most common reasons for readmission in this group were classified as respiratory (1.5% [11/750]), rehabilitation (1.2% [9/750]), infection (0.7% [5/750]), gastrointestinal (0.7% [5/750]), cardiovascular (0.5% [4/750]), and urological (0.5% [4/750]). Thirty-day mortality was 0.3% with standard EVAR and 0.4% with fast-track EVAR.

**Perioperative health care costs**

The total cost to the health care system over 30 days from discharge following standard EVAR was $30,730−$29,300 incurred by hospitals and $1430 incurred by health care payers. In the fast-track EVAR group, the total cost was $26,730−$26,320 incurred by hospitals and $410 incurred...
by health care payers. Ultimately, total health care costs were $4000 lower with fast-track EVAR vs standard EVAR, with $2980 in savings to hospitals and $1030 savings to health care payers. The primary factors contributing to these cost reductions were shorter hospital stay, shorter operating room time, lower rate of intensive care monitoring, and fewer postdischarge MAEs (Table 4). When accounting for the combined uncertainty in all model parameters using Monte Carlo simulations, the mean savings with fast-track EVAR remained $4000, with a 95% confidence interval ranging from $3130 to $4830 (Figure 1).

**Sensitivity analyses**

This study investigated the primary perioperative elements that may substantially influence costs with standard EVAR and fast-track EVAR. However, two elements of the fast-track EVAR program—no general anesthesia and bilateral percutaneous access—were not included in base-case cost models given the absence of these data within Medicare claims. Therefore, we performed sensitivity analyses to assess the potential impact of these exclusions. Regarding the choice of anesthesia, we assumed a $210 cost reduction with regional/local anesthesia vs general anesthesia (Premier Research Services, email communication, November, 2015), and that 81% of the standard EVAR patients underwent general anesthesia. Given that general anesthesia was used in 2.4% of the fast-track EVAR patients, inclusion of anesthesia type in the model would have yielded approximately $170 per patient in additional cost savings with fast-track EVAR. Regarding the choice of vascular access, the cost of vascular closure devices for bilateral PEV AR was estimated at $1050 and the cost of a cutdown tray and sutures was estimated at $390. Assuming 23% of the standard EVAR patients are treated with bilateral PEVAR vs 100% in the fast-track EVAR group, this would yield an additional expense of approximately $510 per patient with fast-track EVAR had vascular closure devices been modeled. Collectively, these data suggest that exclusion of anesthesia and vascular closure devices from the model would decrease the savings with fast-track EVAR relative to the base-case results by approximately 9%—from $4000 to $3660 per patient.

A second sensitivity analysis was performed to estimate the impact associated with the inability to match groups on renal insufficiency and congestive heart failure. For this analysis, we excluded 32 fast-track EVAR patients with renal insufficiency or congestive heart failure. When comparing 218 (87%) fast-track EVAR patients without renal insufficiency or congestive heart failure to standard EVAR patients, total health care cost savings with fast-track EVAR increased by approximately 8%—from $4000 to $4310 per patient.

Ultimately, the base-case cost results in this study ($4000 savings with fast-track EVAR) were robust to modeling assumptions. The additional $340 cost related
to exclusion of anesthesia and vascular access data from the model was balanced by an additional cost savings of $310 when adjusting for the greater frequency of renal insufficiency and congestive heart failure in the fast-track EVAR group.

A final sensitivity analysis evaluated the 216 (86%) patients who completed all elements of the fast-track EVAR protocol (bilateral PEVAR, local/regional anesthesia, no intensive care monitoring, and next-day hospital discharge). In this group, total health care cost savings relative to standard EVAR were $4650, with $3580 in savings to hospitals and $1060 in savings to health care payers (Figure 2).

**Discussion**

The results of this study demonstrate that a fast-track EVAR protocol using a 14 Fr endograft in well-selected patients results in shorter procedure time, lower utilization of intensive care monitoring, faster discharge, lower incidence of MAE, and lower readmission rates compared to a matched sample undergoing standard EVAR. Total perioperative health care costs were $4000 lower with fast-track EVAR, with cost reductions of $2980 to hospitals and $1030 to health care payers. Considering that EVAR without major complication or comorbidity has been reported to result in $4000 in negative operating margins per case,\(^2\)

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Table 3 Perioperative utilization and cost estimates in matched patients undergoing standard and fast-track EVAR\(^a\)

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Utilization</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Standard EVAR</td>
<td>Fast-track EVAR(^b)</td>
</tr>
<tr>
<td>No. of patients</td>
<td>750</td>
<td>250</td>
</tr>
<tr>
<td><strong>Hospital utilization</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Procedure</td>
<td>– (–)</td>
<td>– (–)</td>
</tr>
<tr>
<td>Operating room time (hrs)</td>
<td>2.83±0.13</td>
<td>2.30±0.05(^†)</td>
</tr>
<tr>
<td>Hospital stay (d)</td>
<td>1.69±0.05</td>
<td>1.16±0.07(^†)</td>
</tr>
<tr>
<td>Intensive care monitoring</td>
<td>360 (48.0)</td>
<td>11 (4.4)(^†)</td>
</tr>
<tr>
<td>Major adverse event</td>
<td>12 (1.6)</td>
<td>1 (0.4)</td>
</tr>
<tr>
<td>Secondary intervention</td>
<td>18 (2.4)</td>
<td>0(^*)</td>
</tr>
<tr>
<td><strong>Postdischarge utilization</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major adverse event</td>
<td>54 (7.2)</td>
<td>0(^†)</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>26 (3.5)</td>
<td>0(^†)</td>
</tr>
<tr>
<td>Renal failure</td>
<td>17 (2.3)</td>
<td>0(^†)</td>
</tr>
<tr>
<td>Respiratory failure</td>
<td>8 (1.1)</td>
<td>0</td>
</tr>
<tr>
<td>Stroke</td>
<td>2 (0.3)</td>
<td>0</td>
</tr>
<tr>
<td>Bowel ischemia</td>
<td>1 (0.1)</td>
<td>0</td>
</tr>
<tr>
<td>Paralysis</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Secondary intervention</td>
<td>4 (0.5)</td>
<td>0</td>
</tr>
<tr>
<td>Laparoscopic procedure</td>
<td>3 (0.4)</td>
<td>0</td>
</tr>
<tr>
<td>Endovascular procedure</td>
<td>1 (0.1)</td>
<td>0</td>
</tr>
<tr>
<td>Other readmission(^c)</td>
<td>23 (3.1)</td>
<td>4 (1.6)</td>
</tr>
<tr>
<td>All-cause readmission</td>
<td>51 (6.8)</td>
<td>4 (1.6)(^†)</td>
</tr>
<tr>
<td>Death</td>
<td>2 (0.3)</td>
<td>1 (0.4)</td>
</tr>
<tr>
<td><strong>Model assumptions</strong></td>
<td>Value</td>
<td></td>
</tr>
<tr>
<td>Primary health care payer of postdischarge costs(^d)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medicare</td>
<td>83%</td>
<td></td>
</tr>
<tr>
<td>Private payer</td>
<td>17%</td>
<td></td>
</tr>
<tr>
<td>Private payer to Medicare payment ratio(^e)</td>
<td>1.7 (1.4, 1.9)</td>
<td></td>
</tr>
</tbody>
</table>

Notes: \(^a\)Values are mean ± standard error or count (percent). \(^b\)Comparisons of fast-track EVAR to matched standard EVAR: \(^*\)P<0.05; \(^†\)P<0.01; \(^‡\)P<0.001. \(^c\)Excludes readmission for major adverse event or secondary intervention. \(^d\)Derived from National Inpatient Sample. \(^e\)Estimate represents mode (min, max) from triangular distribution derived from private payer to Medicare payment ratio for cardiovascular procedures. \(^f\)Cost included within other postdischarge utilization categories.

Abbreviation: EVAR, endovascular aneurysm repair.
adoption of a fast-track EVAR program may help to achieve revenue neutrality within this subset of patients.

Several surgical specialties have introduced short-stay protocols for certain procedures with the goal of reducing complication rates, facilitating postoperative recovery, expediting hospital discharge, and reducing early readmissions.\textsuperscript{23,24} Yet few studies have investigated the financial implications of such treatment pathways in EVAR. Moscato et al,\textsuperscript{12} reported a $2400 savings in hospital costs from a retrospective review of patients who required only uncomplicated monitoring with same-day discharge after EVAR. Lachat et al,\textsuperscript{11} reported a $2800 hospital cost savings in matched patients with outpatient EVAR vs inpatient EVAR. Al-Zuhir et al,\textsuperscript{9} reported hospital savings of $2500 in patients undergoing short-stay EVAR with next-day discharge. Importantly, none of these studies were performed in the US and with a requirement of an overnight stay. Since EVAR reimbursement requires inpatient admission at most private institutions in the US, the design of the current study attempted to assess cost savings with fast-track EVAR within the framework of current reimbursement policy within the US. Hospital cost savings of $2980 with fast-track EVAR in the current study were in line with these previous studies,

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Standard EVAR (n=750)</th>
<th>Fast-track EVAR (n=250)</th>
<th>Mean cost savings per patient with Fast-track EVAR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total perioperative costs</td>
<td>30,730</td>
<td>26,730</td>
<td>4000</td>
</tr>
<tr>
<td>Hospital costs</td>
<td>29,300</td>
<td>26,320</td>
<td>2980</td>
</tr>
<tr>
<td>Procedure</td>
<td>24,340</td>
<td>23,390</td>
<td>950</td>
</tr>
<tr>
<td>Ward</td>
<td>4140</td>
<td>2850</td>
<td>1290</td>
</tr>
<tr>
<td>Intensive care monitoring</td>
<td>600</td>
<td>60</td>
<td>550</td>
</tr>
<tr>
<td>Secondary intervention</td>
<td>130</td>
<td>&lt;10</td>
<td>120</td>
</tr>
<tr>
<td>Major adverse event</td>
<td>90</td>
<td>20</td>
<td>70</td>
</tr>
<tr>
<td>Health care payer costs</td>
<td>1430</td>
<td>410</td>
<td>1030</td>
</tr>
<tr>
<td>Other readmission\textsuperscript{a}</td>
<td>520</td>
<td>270</td>
<td>240</td>
</tr>
<tr>
<td>Major adverse event</td>
<td>820</td>
<td>100</td>
<td>730</td>
</tr>
<tr>
<td>Secondary intervention</td>
<td>90</td>
<td>40</td>
<td>50</td>
</tr>
</tbody>
</table>

Note: \textsuperscript{a}Excludes readmission for major adverse event or secondary intervention.

Abbreviation: EVAR, endovascular aneurysm repair.
and were primarily attributable to shorter length of stay, shorter procedure time, and lower intensive care monitoring requirements.

To the author’s knowledge, no fast-track EVAR study has examined costs following hospital discharge. The fast-track EVAR protocol in the current study resulted in a $1030 savings to health care payers over the first 30 days following discharge, largely due to fewer MAEs requiring readmission. Hospital readmission has become the focus of quality improvement efforts owing to the added cost placed on patients and the health care system. In the fast-track, EVAR group, mean hospital stay was 1.2 days and the readmission rate was 1.6%. This in comparison to 1.7 days hospital stay and 6.8% readmission rate with standard EVAR. These results suggest that the hospital benefits of a fast-track EVAR protocol, which includes early discharge, are not offset by higher risk of post-discharge complications and associated readmissions.

Strengths of this research are a large sample of 1000 EVAR patients, patient matching to adjust for group differences in baseline characteristics, determination of utility and cost of EVAR using nationally representative data adjusted for primary health care payer mix, and robustness of conclusions to various sensitivity analysis assumptions. There are several limitations of this study that warrant additional discussion. First, since the fast-track EVAR study followed patients through 1-month, we were unable to determine if clinical benefits and cost savings with fast-track EVAR persisted over longer follow-up. Second, all patients who underwent fast-track EVAR were treated with a 14 Fr endograft and, therefore, these results may not be generalizable to other stent grafts or to other fast-track pathways. While endograft costs account for 38% to 54% of the hospital EVAR costs, endograft cost was not included in this study since stent graft manufacturer data were unavailable in the standard EVAR group and that these costs vary by institution. Regardless, the results of this study are robust to endograft cost assumptions since the 14 Fr endograft for fast-track EVAR remains less costly vs standard EVAR up to $4000 in additional endograft cost. Third, data for the standard EVAR group were derived from Medicare claims. The fast-track and standard EVAR groups were comprised of patients undergoing elective EVAR with comparable demographic and medical history after patient matching. However, despite this matching, other important variables such as aortoiliac morphology were not available within Medicare claims. Such unmeasured variables may have influenced patient outcomes and, thus, are important sources of possible bias. Finally, and perhaps most importantly, fast-track EVAR is only intended for well-selected patients. Patients with heavily calcified or extremely tortuous femoral arteries may be poor candidates for bilateral percutaneous vascular access and patients with major comorbidities may require intensive care monitoring or prolonged hospitalization. While it is plausible that certain elements of the fast-track EVAR protocol could be customized based on individual patient characteristics, the clinical utility of such an individualized strategy has not been studied.

Conclusions
The results of this study demonstrate that a fast-track EVAR protocol using a 14 Fr endograft in well-selected patients results in shorter procedure time, lower utilization of intensive care monitoring, faster discharge, lower incidence of MAE, and lower readmission rates compared to a matched sample undergoing standard EVAR. Further, total perioperative health care costs were $4000 lower with fast-track EVAR, with cost reductions of $2980 to hospitals and $1030 to health care payers. Additional studies with longer follow-up are needed to confirm that fast-track EVAR continues to offer significant clinical and cost benefits.

Data sharing statement
Individual deidentified participant data and study-related documents will not be made available.

Acknowledgment
Endologix, Inc. provided financial support for this study. Endologix, Inc. is the sponsor of the LIFE registry and was involved in study design and study management.

Disclosure
ZK, VGR, EAH, and LEM disclose consultancy with Endologix, Inc. WKN reports personal fees from Trivascular, outside the submitted work. LHA reports personal fees from Miller Scientific Consulting, during the conduct of the study. The authors report no other conflicts of interest in this work.

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