New generation aspiration catheter: Feasibility in the treatment of pulmonary embolism

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AIM: To report our preliminary experience with a new generation aspiration catheter in the treatment of symptomatic pulmonary embolism (PE).

METHODS: A retrospective database search for pulmonary artery embolectomy since introduction of the Pronto .035” and XL extraction catheter (Vascular Solutions, Minneapolis, MN) at our institution in 10/2009 was performed. Ten consecutive patients were identified in which the Pronto .035” or XL catheter was used between 01/2010 and 03/2013. All patients were referred for catheter based embolectomy due to contraindications to systemic lysis, or for being in such a critical clinical condition that immediate percutaneous treatment deemed warranted. The computed tomography (CT) right to left heart ratio as predictor for the severity of the PE was retrospectively evaluated on standard axial views. The difference between pre- and post-procedure pulmonary pressure measures was taken to assess the procedural effect.

RESULTS: Extensive PE was confirmed angiographically in all patients. Measured right- to left ventricle (RV/LV) ratios were elevated beyond one in seven of the eight available CTs. Acute procedural success defined as clinical removal of visible thrombus and improvement in mean pulmonary artery pressure was seen in all recorded patients (n = 8), the mean pulmonary pressures declined from a median (range) of 35.5 (19-46) to 23 (10-37, P = 0.008) mmHg. Neither death nor other complications occurred intra- or immediately periprocedural, yet short term mortality within 30 d was found in 6 out of 9 patients, one patient was lost in follow up. The cause of death within 30 d in the 6 patients was identified as: Circulatory failure in direct connection with the PE (n = 2), stroke, sepsis, or succumbing to malignancy in a hospice setting (n = 2).

CONCLUSION: Success in thrombus removal with improved pulmonary hypertension and systemic hypotension suggests this aspiration technique to be effective. Aspiration catheters should be part of further trials.

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Key words: Pulmonary embolism; Aspiration; Catheter; Thrombectomy; Pulmonary artery

Core tip: We present a new aspiration catheter for use in pulmonary embolism (PE) (Pronto .035” and XL extraction catheter, Vascular Solutions, Minneapolis, MN) in a case series of ten patients. The aspiration catheter allowed fast thrombus removal and lowered mean pulmonary artery pressure. No peri-procedural complications occurred, but high 30-d mortality remained. Catheter based aspiration embolectomy should be considered in acute symptomatic PE, since it is fast and does not require additional special equipment, thus signifying a widely applicable technique. Aspiration embolectomy should be included in further trials treat-
ting symptomatic PE.

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INTRODUCTION

The treatment of massive pulmonary embolism (PE) remains an ongoing challenge due to acute onset of right heart failure and cardiogenic shock limiting the time for therapeutic success. Systemic thrombolysis is often contraindicated and when given involves > 20% complications including 3% intracranial bleeding with a 30 d mortality of 25%-65% depending on severity and case selection[1,2]. In this setting intervention with percutaneous mechanical thrombectomy is appealing, aiming to rapidly reduce the clot burden, pulmonary artery pressure, right heart strain, and to reestablish circulation[3,4].

Several devices have been developed for this task, but each comes with its limitations. The AngioJet (Medrad Interventional/Possis, Minneapolis, MN) and Hydrolyzer (Cordis, Miami, FL) thrombectomy systems use negative pressure created by high-pressure saline injection to aspirate the thrombus[5]. The AngioJet device has been associated with bradyarrhythmias and deaths when used in the pulmonary arteries and carries a black box warning for that use[6,7]. Angioplasty balloons or a rotational pigtail catheter (Cook, Bloomington, IN) have been applied to mechanically fragment a large central thrombus into smaller pieces. However, the fragments can end up occluding previously perfused arteries, potentially increasing pulmonary artery pressures and right heart strain[8]. The principle of aspiration embolectomy has proven successful in small and moderate sized arteries like coronaries. The Greenfield suction embolectomy catheter (Boston Scientific, Natick, MA) was the only PE catheter device approved by the FDA. Bulky design with difficult steerability compromised widespread utilization[9]. Lang et al[10] reported successful treatment with a 14 F 90-cm-long Ultrathane non tapered catheter (Cook, Bloomington, IN), but this catheter was not steerable. Extensive manipulation for repeat access to specific pulmonary artery sites was required for each aspiration pass. This slowed the procedure greatly. Both are no longer available.

We report our preliminary experiences with a new generation of aspiration catheters, the 10 F Pronto .035” and 14 F XL extraction catheter (Vascular Solutions, Minneapolis, MN) in the treatment of acute symptomatic pulmonary embolism. Compared to the preceding suction catheters, these catheters are quickly manipulated over a wire, combining protection from pulmonary artery wall perforation with a rapid return to the target position after a run of aspiration.

MATERIALS AND METHODS

Institutional review board (IRB) approval was obtained. A retrospective database search for pulmonary artery embolectomy since introduction of the catheters in 10/2009 was performed. Ten consecutive patients were identified in which the Pronto .035” or XL catheter were used between 01/2010 and 03/2013.

Patient characteristics are demonstrated in Table 1. The median age of patients was 62.5 (range 33-74) years. All patients were referred for catheter based embolectomy due to contraindications to systemic lysis, or the circulatory status was judged so critical that thrombolytic therapy was deemed not likely to be effective in the available time[1]. The computed tomography (CT) right to left heart ratio as predictor for the severity of the PE was measured on standard axial views[11-13]. Tissue plasminogen activator (tPA) and systemic Heparin were given at the discretion of the operator in dosages as shown in Table 1. Recorded blood loss varied widely from the common 120-300 mL to the maximum volume of 800 mL in one patient. An intraprocedural blood transfusion was required in that patient. Most patients received an IVC filter as secondary prophylaxis after aspiration.

The nonparametric Wilcoxon signed rank test was used to examine the paired difference between pre- and post-procedure pulmonary pressure measures. Given the small sample size, which limited the power for detecting differences, survival was not statistically compared. Two-sided P-values less than 0.05 were considered statistically significant.

Use of the Pronto .035” or XL catheter is FDA approved in peripheral arteries or veins and is off-label in the pulmonary arteries. Embolectomy is achieved through a side-port proximal to the tip of the catheter (Figure 1). A pigtail tip configuration in 8 and 14 F is provided to improve large embolus aspiration in the large diameter of the main pulmonary arteries, while the straight and angled tip configuration enables advancement in segmental and branch arteries. The side-port has radio-opaque marker to allow controlled positioning in clot. Aspiration is achieved via a lockable 60 mL syringe connected to a sideport. Suction control is maintained and modified during use with a roller clamp near the syringe. In 8 out of 10 cases the 10 F Pronto .035” catheter was used, since the 8 and 14 F XL catheter only became available in 01/2012. In the remaining 2 cases the 14 F straight and pigtail catheters were used.

The catheter system was introduced through an appropriately sized 10 or 14 F sheath in the right common femoral vein. Access to the main pulmonary arteries was gained according to operator preferences, (six different attendings), either with a Montefiore pulmonary artery or pigtail catheter (Cook, Bloomington, IN) and pressure measurements were obtained. After angiography and positioning of a stable exchange length working wire in the
target artery, the embolectomy catheter was introduced. XL 14 F pigtail catheters were used with rotation under suction to sweep the entire diameter of the large central arteries. The angulated 10 F Pronto \(0.035"\) catheter and of the main right or left central pulmonary artery during the withdrawal movement. If necessary, additional angiographic catheters were used to select challenging branch arteries. The number of passes and endpoint of the procedure was at the discretion of the operator, with angiographic flow improvement combined with improved pulmonary or systemic pressures as the usual criteria to halt the embolectomy (Figure 2).

**RESULTS**

All patients in our series were confirmed to have extensive PE on angiography. As risk factor for PE prior surgery was identified in 5 cases, malignancy as risk factor for PE and adverse outcome\(^1\) was found in 6 patients. Measured right- to left- ventricle (RV/LV) ratios were elevated beyond one in seven of the eight available CTs (Table 1).

Acute procedural success defined as clinical removal of visible thrombus and improvement in mean pulmonary artery pressure (average of systolic and diastolic pressures) was seen in all recorded patients \(n = 8\), Figure 3; the mean pulmonary pressure declined from a median (range) of 35.5 (19-46) to 23 (10-37, \(P = 0.008\) mmHg. No deaths occurred intra- or immediately periprocedural, also no minor complication such as access hematoma or arrhythmia was observed.

A short term mortality within 30 d was found in 6 out of 9 patients, one patient was lost in follow up. The cause of death within 30 d in the 6 patients was identified as: Circulatory failure in direct connection with the PE \(n = 2\), stroke, sepsis, or succumbing to malignancy in a hospice setting \(n = 2\).

**DISCUSSION**

Consistent success in thrombus removal with improvement in pulmonary hypertension and systemic hypotension suggests this aspiration technique is both successful and of physiologic importance in acute symptomatic PE.

Kucher \(et\ al\)\(^{17}\) defined the requirements for an ideal percutaneous PE thrombectomy catheter as being (1) highly maneuverable, (2) effective in thrombus removal, and (3) safe without damaging cardiac or pulmonary structures. The presented catheter system seems to fulfill all of these requirements, and absence of complex and costly technology should favor widespread use with a real impact on this insidious and often fatal disease. Having two configurations available allowed to remove the most urgent central thrombus burden in the right or left main pulmonary artery first, whereas the angled tip and straight configuration was then used to reestablish flow in obstructed inter- and segmental arteries. The amount of discarded blood in our case series varied widely as well as the concomitant use of thrombolytics. Cell sav-

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**Table 1 Characteristics of 10 patients treated with percutaneous aspiration for pulmonary embolism**

| Patients | Age (yr)/Sex | Pertinent risk factor | Axial CT RV/LV ratio | Systemic pressure pre therapy (Sys/Dia/HR) (mmHg) | Systemic pressure post therapy (Sys/Dia/HR) (mmHg) | Pulmonary pressure pre therapy (Sys/Dia/Mean) (mmHg) | Pulmonary pressure post therapy (Sys/Dia/Mean) (mmHg) | Concomitant Therapeutics | Death 30d |
|----------|--------------|-----------------------|----------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|------------------------|---------|
| Patient 1 | 67 F         | Malignancy, Surgery   | 1.6                  | 90/66/145                                       | 81/65/135                                       | 53/37/46                                       | 48/31/37                                       | TPA (mg)               | 6/6000   |
| Patient 2 | 74 M         | MVA, Surgery          | 1.7                  | 120/79/133                                      | 148/102/122                                     | 53/23/33                                       | 32/13/19                                       | Heparin Bolus (U)      | N       |
| Patient 3 | 33 F         | Neurosurgery          | 1.6                  | 100/76/116                                      | 94/51/124                                       | 54/27/38                                       | 11/27/1934                                    | N                      | Y       |
| Patient 4 | 47 M         | MVA, Surgery          | 1.3                  | 133/57/143                                      | 115/54/131                                      | 7/19/1943                                      | 4/14/1931                                      | Y                      |         |
| Patient 5 | 47 F         | Malignancy, PEs, COPD | 0.8                  | 88/65/45                                        | 87/52/37                                        | NA/NA/24                                       | NA/NA/10                                       | N                      |         |
| Patient 6 | 57 M         | Malignancy            | NA                   | 126/102/112                                     | 102/79/97                                       | NA/35/40                                       | NA                                            | 0                      | N       |
| Patient 7 | 64 M         | Malignancy, Surgery, COPD | 1.3             | 103/76/115                                      | 119/94/90                                       | 60/20/40                                       | NA                                            | 0                      | N       |
| Patient 8 | 62 M         | Diabetes, Hypertension| 1.4                  | 111/74/80                                      | 104/80/70                                       | 58/27/38                                       | 44/18/31                                      | 16/5000                | N       |
| Patient 9 | 73 M         | Malignancy            | 1.5                  | 116/72/115                                      | 116/69/131                                      | NA/NA/30                                       | NA/NA/27                                       | 6                      | Y       |
| Patient 10| 63 M         | Malignancy            | NA                   | 165/82/108                                      | 66/57/62                                        | 39/15/25                                       | 25/14/18                                      | 4.5                    | Y       |

M: Male; F: Female; COPD: Chronic obstructive pulmonary disease; Dia: Diastolic; HR: Heart Rate; NA: Not available; PEs: Prior pulmonary embolism; RV/LV ratios: Right to left ventricle ratio; Sys: Systolic; tPA: Tissue plasminogen activator; CT: Computed tomography.
Ing techniques are routine in operative cases, and may well be appropriate here. Requirements for blood trans-

fusions need to be part of future investigations as well as establishing standardized concomitant drug regimens

and firm therapeutic endpoints.

It is evident that a small retrospective series like ours has considerable limitations pertaining to generalization of our findings. It is also possible that a variety of treatment strategies may have their roles, since different types of thrombi and emboli may be encountered and one treatment method may not fit all.

Although mortality has been reported to range from about 20%-65% in those presenting with shock, the high 66% mortality within 30 d (Table 1) could bring into question the efficacy of the aspiration system. Taking the RV/LV ratio of one as indicator for severity of the PE, seven of our measured eight patients presented with largely elevated ratios from 1.3-1.7. Even though the RV/LV ratio is a controversial prognostic indicator, it seems to outperform other radiologic surrogate measurements. Similarly, six patients had underlying malignancy as risk factor for adverse short term outcome. The high mortality appears therefore rather to be a reflection of the critical condition of the patients than to refute the efficacy of percutaneous embolectomy. No death could be linked to a procedural complication.

Considering that conservative management with systemic tPA results in numerous complications and death rates up to 65%, we propose an earlier role for catheter based embolectomy than in a moribund state.

To generate best-possible evidence in the treatment of massive pulmonary embolism we suggest that percutaneous aspiration catheters be part of further trials and

Figure 2  Treatment of massive pulmonary embolism with an aspiration catheter. A: Axial contrast enhanced computed tomography (CT) at time of diagnosis: Extensive bilateral PE with complete occlusion of the right pulmonary artery with thrombus in the ascending branch and the inferior trunk, partial obstruction of the left upper lobe branch (arrows). Left subcutaneous emphysema from vigorous resuscitation is noted; B: Digital Subtraction Angiography of the right pulmonary artery (RAO): Angiographic findings confirm complete obstruction of the ascending branch (diamond) and the interlobar artery of the right pulmonary artery (arrow); C: Angiographic imaging of pulmonary artery embolus aspiration. A 10 F aspiration catheter (Pronto .035”) is advanced into the thrombus over a guidewire (arrow). Radiopaque marking of the catheter tip enables controlled positioning of the aspiration side hole just proximal to the tip in the thrombus (Triangle); D: Digital Subtraction Angiography of the right pulmonary artery (RAO): After aspiration maneuvers, selective angiography in the interlobar artery confirms reestablished contrast flow in the interlobar artery with remaining thrombus (arrow), reflux in the ascending branch (diamond) is visualized.

Figure 3  Mean pulmonary pressure (mmHg) pre- and post-procedure (n = 8) note: Dots indicate median values. Photoshop (Adobe) was used for imaging formatting.

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be included in nationwide pulmonary embolism treatment registries.

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