EP63 THE TAPAS STUDY: THROMBECTOMY ASPIRATION POST-MARKET-STUDY IN ACUTE STROKE

Department of Neuroradiology, Hospital Universitario 12 de Octubre, Madrid, Spain.
Hospital Virgen de la Arrixaca, Murcia; Centro Hospitalario Universitario Donostia, San Sebastian; Hospital Universitario de Santiago de Compostela, Santiago de Compostela, Spain

Introduction The Q Aspiration catheter design has shown increased aspiration force in bench studies. The objective of this first clinical study was to assess safety and performance of the Q™ Catheter in a real-world setting. The study was a multi-center, observational, post market study utilizing the Q Catheter as first-line therapy during mechanical thrombectomy in Spain.

Methods Patients with AIS treated with Q as first line therapy between March 2019 and January 2020 were identified in four centers in Spain. Inclusion criteria included 18–85 years of age, anterior or posterior LVO, ASPECTs 6–10 and within 8 hours of onset/last known well. Baseline demographics, procedural data, post procedure neuroimaging, and clinical outcomes were reported. A follow up visit was conducted to assess mRS.

Results There were 45 patients included. Average age at presentation was 72.4 and 53.3% were male. Thrombolytic therapy was given in 46.7% (21/45) and NIHSS was 14.4 (range 1–26). Final mTICI 2b-3 was achieved in 93% of patients (42/45). First pass success of mTICI 2b-3 with only a Q Catheter was achieved in 49% (21/43), increasing to 56% (24/43) when a stent retriever was used as an anchor. Overall successful revascularization using the Q Catheter was 77% (33/43). ENT was 2% (1), sICH rate was 2% (1), and mortality during the study period was 13% (6). At follow-up, 55.5% of subjects had a mRS of 0–2.

Conclusion The Q Catheter showed good deliverability and revascularization rates in this real-world setting with no safety concerns.

REFERENCE

Disclosure Consultant MIVI Neuroscience, Inc.

EP64 SYSTEMATIC REVIEW AND ANALYSIS OF PRE-ClinICAL SIDE-BY-SIDE COMPARISONS OF EMBOLTRAP VERSUS SOLITAIRE PERFORMANCE

M Mirza, R McCarthy, M Gilvary. 1Neuro Thromboembolic Initiative, 2Cerovens, Galway, Ireland

Introduction The first pass effect (FPE) has gained acceptance as the new measure of success for mechanical thrombectomy. Lacking clinical trials that compare devices, in-vitro studies may be useful to investigate the impact of device designs to FPE.

Aim of the Study To systematically compare EmboTrap performance using side-by-side generated in-vitro comparative studies

Methods A systematic review of literature for pre-clinical data containing in-vitro thrombectomies was performed in Pubmed. Articles were included if multiple devices including EmboTrap were used in the same models, techniques, and blood clot analogs. Individualized data were extracted, and a pooled analysis was performed. Comparisons of recanalization success between stent-retrievers and different clot types were performed using two-way analysis of variance (ANOVA) and a two-sample t-test with Tukey’s honestly significant.

Results An initial search revealed 277 articles, 4 articles met the inclusion criteria. Only EmboTrap and Solitaire devices were consistently used in all 4 articles, limiting comparisons to only these two devices. Three articles described outcomes as first-pass complete clot removal, while one article allowed up to 3 passes, for a total of 210 thrombectomy attempts. Of the three clot types studied, friable and tough clots were recalanized less effectively than standard clots (p<0.01), but not different from each other (p>0.05). EmboTrap had better recanalization outcomes than Solitaire (p<0.01), particularly observed for friable (p<0.05) and standard (p<0.05) clots, but not for tough clots (p>0.05).

Conclusions Pre-clinical data, particularly from combined independent studies, can provide insightful comparisons between device designs.

REFERENCE

Disclosure MM, RM, and MG are employees of Cerovens.
had left side ACM occluded, while female patients had the left ACM occlusion, with a sudden onset of right/left hemiparesis, right gaze preference and aphasia.

The first case (58 years male) we performed mechanical thrombectomy with aspiration. CT angiography revealed acute left ACM occlusion, 15 mm M1 thrombus in the left ACM, NIHSS score 13 before the procedure (19.45 hours) and after the procedure 7 points NIHSS (00.45 hours). All patients underwent MRI and MRA examination next day was performed presenting patent recanalized arteries.

**Intervention** Following confirmation of lesion amenable to mechanical thrombectomy, patients underwent successful complete reperfusion TICI (3) revascularization of ACM territories, four of which using the stent retriever.

**Conclusion** Endovascular treatment has proven to be safe and efficacious for AIS. Our patients recovered and were discharged within 7–11 days after ictus.mRS after 90 days 0 at two (male) patients, mRS 1 at two patients and one patient is mRS 3.

These case series demonstrate the clinical efficacy, safety, and favorable clinical outcome of first mechanical thrombectomies performed in Kosovo.

**REFERENCES**


**Disclosure** Nothing to disclose