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Corresponding author
Dr Alan C Cameron BSc (Hons), MB ChB, MRCP
Institute of Cardiovascular and Medical Sciences
University of Glasgow, 126 University Place
Glasgow, United Kingdom, G12 8TA
Tel: +44 (0)141 330 8271
Email: alan.cameron.2@glasgow.ac.uk

PROFESSIONAL GUIDELINE VERSUS PRODUCT LABEL SELECTION FOR TREATMENT WITH IV THROMBOLYSIS: AN ANALYSIS FROM SITS REGISTRY

Authors
Alan C Cameron¹, James Bogie¹, Azmil H Abdul-Rahim², Niaz Ahmed³, Michael Mazya³, Robert Mikulik⁴, Werner Hacke⁵, Kennedy R Lees¹, for the Safe Implementation of Treatments in Stroke (SITS) Investigators

¹ Institute of Cardiovascular and Medical Sciences, University of Glasgow, United Kingdom.
² Institute of Neuroscience and Psychology, University of Glasgow, United Kingdom.
³ Department of Neurology, Karolinska University Hospital and Department of Clinical Neuroscience, Karolinska Institute, Stockholm, Sweden.
⁴ International Clinical Research Centre, Neurology Department, St Anne’s Hospital and Masaryk University, Brno, Czech Republic.
⁵ Department of Neurology, University of Heidelberg, Germany.
ABSTRACT

Background
Thrombolysis usage in ischaemic stroke varies across sites. Divergent advice from professional guidelines and product labels may contribute.

Methods
We analysed SITS-International registry patients enrolled January 2010 through June 2016. We grouped sites into organisational tertiles by number of patients arriving ≤2.5h and treated ≤3h, percentage arriving ≤2.5h and treated ≤3h, and numbers treated ≤3h. We assigned scores of 1-3 (lower/middle/upper) per variable and 2 for on-site thrombectomy. We classified sites as lower-efficiency (summed scores 3-5), medium-efficiency (6-8) or higher-efficiency (9-11).

Sites were also grouped by adherence with European product label and ESO guideline: “label adherent” (>95% on-label), “guideline adherent” (≥5% off-label, ≥95% on-guideline) or “guideline non-adherent” (>5% off-guideline). We cross-tabulated site-efficiency and adherence. We estimated the potential benefit of universally selecting by ESO guidance, using onset-to-treatment time-specific numbers needed to treat for day 90 mRS 0-1.

Results
56,689 patients at 597 sites were included: 163 sites were higher-efficiency, 204 medium-efficiency and 230 lower-efficiency. 56 sites were “label adherent”, 204 “guideline adherent” and 337 “guideline non-adherent”. There were strong associations between site-efficiency and adherence (P<0.001). Almost all “label adherent” sites (55, 98%) were lower-efficiency.

If all patients were treated by ESO guidelines, an additional 17,031 would receive alteplase, which translates into 1,922 more patients with favourable 3-month outcomes.
Conclusion

Adherence with product labels is highest in lower-efficiciency sites. Closer alignment with professional guidelines would increase patients treated and favourable outcomes. Product labels should be revised to allow treatment of patients ≤4.5 hours from onset and aged ≥80 years.
**Background**

Thrombolysis with intravenous recombinant tissue plasminogen activator (IV rt-PA) (alteplase; Actulyse or Activase) is effective and safe for patients with acute ischaemic stroke, yet only a fraction of patients receive treatment\(^1\)\(^-\)\(^9\). The product labels for IV rt-PA in Europe (EU) and the United States (US) are derived from early randomised controlled trials, which excluded important groups\(^10\). The EU label restricts treatment to patients under 80 years, whilst the US label excludes patients greater than 3 hours from symptom onset. IV rt-PA is effective and safe within 4.5 hours of symptom onset\(^11\)\(^-\)\(^13\) and there is clear treatment benefit in the elderly\(^9\),\(^14\)\(^-\)\(^18\).

Professional guidance from the European Stroke Organisation (ESO) and American Stroke Association (ASA) better reflects the evidence base for alteplase\(^19\)\(^-\)\(^21\). The ESO recommend treatment within 4.5 hours with no age limit, whilst ASA guidance in 2013 excluded patients aged >80 years beyond 3 hours, although an update in 2016 acknowledged alteplase is effective within 4.5 hours in the elderly\(^9\),\(^19\)\(^-\)\(^21\). Thus, patients are often treated off-label\(^22\), although this practice is not permitted in many countries and the current product labels therefore restrict the number of patients that can be treated\(^1\),\(^22\).

We aimed to assess variation in the use of IV rt-PA within the Safe Implementation of Thrombolysis in Stroke International Stroke Thrombolysis Registry (SITS-ISTR), in relation to the principal criteria that differ between regional product labels and professional guidelines. Our objective was to assess whether centres’ expertise, measured in terms of efficient patient throughput and treatment logistics, is associated with closer adherence to the EU / US drug labels and professional guidelines; and to estimate the potential impact on treatment rates and clinical outcomes if there were greater alignment of the product labels and professional guidelines. We hypothesised that centres which achieve excellent treatment logistics will adhere more closely with professional guidelines rather than strictly observing the product label for IV alteplase.
Methods

We conducted a retrospective analysis on individual patient data obtained from the SITS-ISTR between January 2010 and June 2016 (Figure 1). SITS-ISTR is a multinational open registry of patients with acute ischaemic stroke who received IV rt-PA. Patients from 597 participating centres were included who had complete information on treating hospital and country, age, gender, onset-to-treatment time (OTT), total National Institutes of Health Stroke Scale (NIHSS), history of diabetes and history of prior stroke. We excluded patients on direct oral anticoagulants or with OTT recorded as >6 hours. Baseline characteristics included data on pre-stroke modified Rankin Scale (mRS), medical history and medications. We also gathered data for on-site use of thrombectomy.

We grouped sites according to their “selection adherence” of alteplase use: label adherent (>95% of patients treated within label), guideline adherent (≥5% of patients treated off-label but ≥95% of patients treated within guideline) or guideline non-adherent (>5% of patients treated off-guideline). We assessed site quality using a tertile based scoring algorithm. Sites were grouped into tertiles according to (i) the volume of patients arriving within 2.5 hours and treated within 3 hours, (ii) the percentage of patients arriving within 2.5 hours and treated within 3 hours, and (iii) the volume of patients treated within 3 hours of stroke onset. We assigned sites a score for each variable: 1 point if the site was within the lower tertile, 2 points for the middle tertile or 3 points for the upper tertile. An additional 2 points were allocated for on-site use of thrombectomy. This resulted in a total score between 3 and 11 for each site. We classified sites with scores of 3 to 5 as ‘lower efficiency’, 6 to 8 as ‘medium efficiency’ and 9 to 11 as ‘higher efficiency’. We tested associations between site efficiency and selection adherence of alteplase use by cross-tabulation and Chi-squared analyses performed in SPSS version 22.0, with a significance level of 5%.

We estimated the potential for clinical benefit if treatment of all patients within our cohort was by the professional guideline versus product label for alteplase. We performed this analysis by applying guideline criteria for treatment with IV rtPA to our entire cohort, and compared this to the
number of patients that would have been treated if the product label criteria were applied. We conducted separate analyses for both European (ESO guideline and EU label) and American (AHA guideline and FDA label) criteria applied to the entire dataset. We calculated the number of patients for whom treatment would have been contraindicated by the product label but recommended by professional guideline. We stratified such patients according to OTT: within 90 minutes, 91 to 180 minutes or 181 to 270 minutes. We used OTT-specific numbers needed to treat (NNT) for a day 90 modified Rankin Scale (mRS) of 0-1 to estimate the number of additional patients that would achieve a favourable outcome if treatment was universally by professional guideline rather than product label (NNT of 4.5 if OTT was 0 to 90 minutes, NNT of 9.0 if OTT 91 to 180 minutes and NNT of 14.1 if OTT 181 to 270 minutes). We divided the number of additional patients that would be treated within each time window by the corresponding NNT to estimate the number of patients that would achieve a favourable outcome.

**Results**

We analysed data from 56,689 patients treated at 597 sites during the study period. Baseline characteristics of the patients are shown in table 1. By our predefined criteria, 163 sites (27%) were classified as ‘higher efficiency’, 204 sites (34%) as ‘medium efficiency’ and 230 sites (39%) as ‘lower efficiency’.

When analysing selection adherence across all sites by the EU product label and ESO guideline, we found that 56 sites (9%) were label adherent, 204 sites (34%) were guideline adherent and 337 sites (56%) were guideline non-adherent. Site efficiency was strongly associated with selection adherence by European criteria (p<0.001) (Figure 2). Among the 56 label adherent sites, 55 (98%) were lower efficiency and one only (2%) was medium efficiency. Of the 204 guideline adherent sites, 92 (45%) were lower efficiency, 75 (37%) were medium efficiency and 37 (18%) were higher efficiency. Among the 337 guideline non-adherent sites, 126 (37%) were higher efficiency, 128 (38%) were medium efficiency and 83 (25%) were lower efficiency. When we judged use in our mainly European dataset against US product label and ASA guideline
criteria, a similar pattern emerged except that guideline non-adherence rose (see online appendix).

IV rt-PA was administered to 5770 patients (10%) beyond European guideline recommendations. This was due to patients treated with a BP greater than guideline recommendations in 4618 patients (8%), an OTT greater than 4.5 hours in 1047 patients (2%) and a combination of elevated BP with an OTT greater than 4.5 hours in 105 patients (0.2%). Among the 5770 patients administered IV rt-PA beyond European guideline recommendations, 3845 (67%) were treated in a higher efficiency site, 1644 (28%) in a medium efficiency site and 281 (5%) in a lower efficiency site (Figure 3).

Within our cohort, 50,919 patients (90%) would receive thrombolysis if treatment was universally delivered by the ESO guideline, compared to 33,888 patients (60%) by the European product label. Thus, an additional 17,031 patients (30%) would receive thrombolysis if treatment was universally delivered according to ESO guidance. This translates into 1,922 patients who would achieve a favourable outcome when measured by OTT-specific NNT for a day 90 mRS of 0-1 (Figure 4).

Discussion

We have demonstrated that strict adherence with the product label for IV rt-PA is greatest in sites that treat lower volumes of patients, have fewer facilities or achieve less impressive in-hospital timelines. Strict adherence with the product label restricts use of IV rt-PA, reducing the number of patients that can be treated and, by implication, that may achieve favourable outcomes. If treatment decisions within our cohort were based upon ESO guidelines rather than the European drug label, an additional 2620 patients would be treated annually across the 6.5 years studied. This translates into an additional 296 patients each year with favourable outcomes. Evidence supporting the selection criteria described in the ESO and ASA professional guidelines is robust and the conclusions of these organisations agree on all major points19-21. The drug product labels
for alteplase require review in both Europe and America, to reflect evidence highlighting the efficacy and safety of IV rt-PA in circumstances that were originally considered contraindications for thrombolysis\textsuperscript{1,24,25}. The key issue is that these labels, which simply control marketing activities and not prescribing per se, should permit the manufacturers to discuss and educate clinicians on the safe treatment of patients within 4.5 hours of symptom onset or aged over 80 years. Revising the European and American product labels to this effect would deliver clinical outcomes consistent with those obtained when treating within the current drug labels, with no adverse effect on mortality\textsuperscript{26}. Alignment of educational messages is desirable and should be conveyed amongst the medical community\textsuperscript{9}.

Our data demonstrate that less efficient sites have the lowest rates of treatment with alteplase off-label, which may in part be attributable to less developed regions being unable to treat off-label\textsuperscript{27}. This is consistent with findings from a previous study using SITS-ISTR data, which demonstrated that higher volume centres have the greatest rates of treatment with alteplase off-label\textsuperscript{28}. Improving the quality of treatment for every patient with acute stroke is a priority of the ESO and World Stroke Organisation (WSO), with the Angels Initiative recently introduced to help achieve this goal. Education of clinicians and revision of the product labels for alteplase will help our effort to deliver excellent care for patients with acute ischaemic stroke worldwide.

It is concerning that we observed high rates of treatment with alteplase beyond professional guidelines. Off-guideline treatment was administered to 10\% of patients by European criteria, which was driven by treatment above BP recommendations and beyond 4.5 hours. Treatment with alteplase off-guideline exposes patients to an increased risk of mortality that is not offset by potential for clinic benefit\textsuperscript{26} and clinicians should avoid this practice. Violations of pre-treatment BP parameters are associated with an increased risk of bleeding and BP should be controlled before treatment with IV rt-PA to reduce the risk of symptomatic intracerebral haemorrhage\textsuperscript{24,25}. Most off-guideline treatment was in higher efficiency sites and programmes discouraging this approach should include all the stroke community.
We designed a measure of site efficiency that acts as a marker of site quality. We allocated points for efficient treatment logistics, the volume and proportion of patients treated promptly and on-site use of thrombectomy. Our aim was to stratify sites according to treatment logistics, patient volume and delivery of comprehensive acute stroke care. Various indicators can be used to assess quality of acute stroke unit care, although not all of these data are available within SITS-ISTR. Our measure of site quality is arbitrary and uses objective information available within SITS-ISTR defined before we accessed the data, which is thus a weakness of our study. The criterion for site quality includes measures derived mainly from OTT and volume of patients, which may disadvantage centres with longer out-of-hospital transportation logistics and smaller sites. We defined BP based upon that recorded at baseline within the SITS registry and cannot be certain that BP was not lowered prior to thrombolysis, which is a limitation.

A further limitation is the retrospective and observational design, although the large volume and accuracy of data collected within SITS-ISTR allows for robust statistical analyses. SITS-ISTR is a predominantly European cohort which is important when considering the generalisability of our findings. Patients managed outside Europe are often in countries with less experienced centres and our results are relevant to these regions. Finally, SITS-ISTR includes patients voluntarily registered by participating centres which could contribute to selection bias, although data from SITS are robust and have been used in similar studies.

Conclusion
We confirmed that strict adherence with the more restrictive product label for alteplase was concentrated among the least active or efficient hospitals, whereas more experienced sites offer treatment based on according professional guideline criteria. However, we found that the busiest and most efficient sites are treating beyond even the professional guidelines, potentially exposing these patients to a risk of increased mortality that is not offset by potential for clinical benefit. We conclude that review and alignment of the marketing approvals for alteplase in acute ischaemic
stroke with the current recommendations of the professional guidelines, to allow treatment of patients ≤4.5 hours from onset and aged ≥80 years, should be coupled with enhanced education to operate within those guidelines to maximise the population safety and effectiveness of thrombolysis for stroke.

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References


26. Lees KR. On-label versus off-label outcomes with intravenous alteplase. 3rd European Stroke Organisation Conference. May 2017; Prague, Czech Republic.


DECLARATIONS

Conflicting interests
KRL is a member of the Stroke Thrombolysis Trialists’ Collaboration that has published pooled individual patient data analyses of the effects of rt-PA in acute ischaemic stroke; a member of the Scientific Committee of SITS; is Past President of the European Stroke Organisation that publishes guidelines on stroke management and coordinates the ESO-Angels project; and has received fees and expenses for data monitoring committees from Boehringer Ingelheim. WH reports no current conflicts. He declares that he served at the SCs of the ECASS 1-4 trials and has been compensated for work in the SCs and for lectures in the past. NA is the Vice Chairman and MM is a Researcher at SITS International, which receives an unrestricted grant from Boehringer Ingelheim for the SITS-International Stroke Thrombolysis Register. ACC, AHAR and JB report no conflicting interests.

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Ethical approval

Ethical approval for this study was obtained from the Safe Implementation of Treatments in Stroke (SITS) Scientific Committee.

Informed consent

Not applicable.

Guarantor

KRL

Contributorship

KRL, ACC, JB and AHAR researched literature and conceived the study. KRL, ACC, JB and AHAR were involved in protocol development, gaining SITS Scientific Committee approval and data analysis. ACC, JB and AHAR wrote the first draft of the manuscript. All authors reviewed and edited the manuscript and approved the final version of the manuscript.

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Selection process of the study population.

### SITS-ISTR
Safe Implementation of Thrombolysis in Stroke
International Stroke Thrombolysis Registry

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>DOAC</td>
<td>Direct Oral Anti-Coagulant</td>
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<tr>
<td>NIHSS</td>
<td>National Institutes of Health Stroke Scale</td>
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<td>OTT</td>
<td>Onset-to-Treatment Time</td>
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Figure 2

Site efficiency and selection adherence with EU Product Label and ESO Guideline.

Key
- Guideline Non-Adherent (>5% off guideline)
- Guideline Adherent (≥5% off label, ≥95% on guideline)
- Label Adherent (>95% on label)
Figure 3

The percentages of patients treated off-guideline grouped by site efficiency, according to ESO guideline criteria (n=5,770).
The percentage of patients that would be treated if the decision was based on ESO guidance versus EU product label: an additional 17,031 patients (30%) would be treated if the decision was based upon ESO guidance, which translates into 1,922 patients achieving favourable outcomes when estimated using OTT-specific NNT for a day 90 mRS of 0-1.

Key

- Red: Treatment with IV alteplase NOT recommended
- Blue: Treatment with IV alteplase recommended
Table 1
Baseline characteristics of the cohort

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<tr>
<td></td>
<td></td>
<td>N=56689</td>
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<tr>
<td>Age (years)</td>
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<tr>
<td></td>
<td>(SD)</td>
<td>(13.1)</td>
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<tr>
<td>Sex (male)</td>
<td>n</td>
<td>30969</td>
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<tr>
<td></td>
<td>(%)</td>
<td>(55%)</td>
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<tr>
<td>Baseline NIHSS</td>
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<td></td>
<td>(IQR)</td>
<td>(6-16)</td>
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<tr>
<td>Onset to treatment time (minutes)</td>
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<tr>
<td></td>
<td>(IQR)</td>
<td>(118-195)</td>
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<td>Atrial fibrillation</td>
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<tr>
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<td>(%)</td>
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<tr>
<td></td>
<td>(%)</td>
<td>(17%)</td>
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<tr>
<td>Previous stroke or TIA</td>
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<td>9067</td>
</tr>
<tr>
<td></td>
<td>(%)</td>
<td>(16%)</td>
</tr>
</tbody>
</table>
IQR indicates interquartile range; NIHSS, National Institutes of Health Stroke Scale
Appendix Figure 1

Site efficiency and selection adherence with American Product Label and ASA guideline.

Key
- Guideline Non-Adherent (>5% off guideline)
- Guideline Adherent (≥5% off label, ≥95% on guideline)
- Label Adherent (>95% on label)

p<0.001
Appendix Figure 2

The percentage of patients treated off-guideline grouped by site efficiency, according to ASA guideline criteria (n=10,690).
Appendix Figure 3

The percentage of patients that would be treated if the decision was based on ASA guidance versus the American product label: there were 13,070 patients (23%) for whom treatment would have been recommended by ASA guidance but not the American product label. Amongst these 13,070 patients, 1,114 would achieve a favourable outcome when estimated by OTT-specific NNT for a day 90 mRS of 0-1.

Key

- Treatment with IV alteplase NOT recommended
- Treatment with IV alteplase recommended