Thrombolysis in very elderly people: controlled comparison of SITS International Stroke Thrombolysis Registry and Virtual International Stroke Trials Archive

Nishant K Mishra, doctoral candidate,1 Niaz Ahmed, neurologist,2 Grethe Andersen, neurologist,3 José A Egido, neurologist,4 Perttu J Lindsberg, professor of applied neurology,5 Peter A Ringleb, neurologist,6 Nils G Wahlgren, professor of neurology,2 Kennedy R Lees, professor of cerebrovascular medicine1 for the VISTA and SITS collaborators

ABSTRACT
Objective To assess effect of age on response to alteplase in acute ischaemic stroke.
Design Adjusted controlled comparison of outcomes between non-randomised patients who did or did not undergo thrombolysis. Analysis used Cochran-Mantel-Haenszel test and proportional odds logistic regression analysis.
Setting Collaboration between International Stroke Thrombolysis Registry (SITS-ISTR) and Virtual International Stroke Trials Archive (VISTA).
Participants 23 334 patients from SITS-ISTR (December 2002 to November 2009) who underwent thrombolysis and 6166 from VISTA neuroprotection trials (1998-2007) who did not undergo thrombolysis (as controls). Of the 29 500 patients (3472 aged >80 (elderly), mean 84.6), data on 272 patients were missing for baseline National Institutes of Health stroke severity score, leaving 29 228 patients for analysis adjusted for age and baseline severity.
Main outcome measures Functional outcomes at 90 days measured by score on modified Rankin scale.
Results Median severity at baseline was the same for patients who underwent thrombolysis and controls (odds ratio 1.6, 95% confidence interval 1.5 to 1.7; Cochran-Mantel-Haenszel P=0.001). The association occurred independently among patients aged ≤80 (1.6, 1.5 to 1.7; P=0.001; n=25 789) and in those aged >80 (1.4, 1.3 to 1.6; P=0.001; n=3439). Odds ratios were consistent across all 10 year age ranges above 30, and benefit was significant from age 41 to 90; dichotomised outcomes (score on modified Rankin scale 0-1 v 2-6; 0-2 v 3-6; and 6 (death) v rest) were consistent with the results of the ordinal analysis.
Conclusions Outcome in patients with acute ischaemic stroke is significantly better in those who undergo thrombolysis compared with those who do not. Increasing age is associated with poorer outcome but the association between thrombolysis treatment and improved outcome is maintained in very elderly people. Age alone should not be a barrier to treatment.

INTRODUCTION
Thrombolysis for acute ischaemic stroke has proved benefits,1-5 but data from randomised trials in patients aged over 80 are limited. About 30% of acute stroke occurs in people aged over 80.5-8 The NINDS (National Institute of Neurological Disorders and Stroke) trial initially restricted enrolment to patients aged up to 80.9 The age criterion was lifted after they enrolled 188 patients in part A of the trial, but they enrolled only 42 very elderly patients.2 All ECASS (European Cooperative Acute Stroke Study) trials applied an upper age limit of 80,3,10 and recent studies with desmoteplase (a fibrin specific plasminogen activator)11,12 also excluded elderly patients. Thus, up to now, the European Medicines Evaluation Agency has not approved thrombolysis with alteplase among patients aged above 80.6 Many experienced centres treat elderly patients but others observe the terms of product approval.13-16

The main reasons for withholding treatment from very elderly patients in clinical practice are fears that advancing age is associated with poorer prognosis with greater risk for haemorrhage and in hospital mortality.17-20 Conversely, a meta-analysis of pooled thrombolysis data concluded that the risks of symptomatic intracerebral haemorrhage did not increase among elderly patients, despite less favourable outcomes.15,20 Less favourable outcomes are expected to occur in elderly patients, mostly because of comorbidity.19,21

The proportion of older people is rising in our society,5,8 and the proportion of those undergoing thrombolysis will decline in the future if patients aged over 80 are not treated.22 In the United Kingdom alone,
the population aged over 80 has doubled since 1982,23 and life expectancy has risen in the rest of Europe and in other countries.24 Effective treatments, however, should not be withheld from older people in the absence of compelling data suggesting unacceptable risk or proved lack of benefit. We hypothesised that absence of compelling data suggesting unacceptable should not be withheld from older people in the Archive, VISTA (www.vista.gla.ac.uk).26

We collated the data of stroke patients who underwent thrombolysis through the SITS-ISTR (Safe Implementation of Treatment in Stroke-International Stroke Thrombolysis Registry, www.sitsinternational.org, held at Karolinska University Hospital, Stockholm) from December 2002 to November 2009. Our control group comprised untreated stroke patients from neuroprotection trials conducted from 1998 to 2007 and held within the Virtual International Stroke Trials Archive, VISTA (www.vista.gla.ac.uk).26

The SITS-ISTR is an ongoing internet based, academic driven, interactive thrombolysis register. The methods of the register, including the procedure for data collection and management, identification of patients, and verification of source data, has been described previously.27 28 In brief, it is a prospective open multinational observational monitoring registry for clinical centres using thrombolysis and other interventions for the treatment of acute ischaemic stroke. The registry is open to all countries, and collects data on patients who receive thrombolytic therapy for acute ischaemic stroke.28

VISTA is a collaborative, not-for-profit, register of stroke trials.26 The treatments studied in these trials range from putative neuroprotectants through anticoagulants and thrombolytic agents to simple rehabilitation measures.26 Unfortunately, the effects of the neuroprotectants have largely been indistinguishable from placebo.29 All trials in VISTA hold necessary review board and regulatory approvals, and all patients agreed to participation. The archive holds only anonymised data, and the source trial is not disclosed as per VISTA guidelines.26 For this analysis, we sought data from the archive from a group of trials in which the investigated drug was a putative neuroprotectant that was neither vasoactive nor interfered with clotting or from placebo groups. Hence, the “controls” were patients who did not receive alteplase in neuroprotection trials but received either placebo or a neuroprotective drug as per the randomisation protocol for each contributing trial. From both of these sources, we collated the demographics, clinical data, and information of functional outcome as measured by the score on the modified Rankin scale after 90 days.

METHODS
Data source and patients
We collated the data of stroke patients who underwent thrombolysis through the SITS-ISTR (Safe Implementation of Treatment in Stroke-International Stroke Thrombolysis Registry, www.sitsinternational.org, held at Karolinska University Hospital, Stockholm) from December 2002 to November 2009. Our control group comprised untreated stroke patients from neuroprotection trials conducted from 1998 to 2007 and held within the Virtual International Stroke Trials Archive, VISTA (www.vista.gla.ac.uk).26

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Patient sample
We downloaded data on 28 136 patients registered in SITS-ISTR from 25 December 2002 to 2 November 2009 and extracted information on 23 336 who had completed 90 day follow-up. Two patients were excluded because of inexact information on age, leaving 23 334. Participating centres agreed to record details of every patient treated with intravenous alteplase at that centre and to enter baseline data as soon as possible after treatment, adding outcomes as they occurred.27

We collated data on 9665 patients from VISTA. Of these, 6371 patients who had ischaemic stroke and did not undergo thrombolysis were selected as controls. We excluded 205 patients with no information on functional outcomes at 90 days, leaving 6166 for analysis. Of the 29 500 patients (n=3472 aged >80), data on baseline National Institutes of Health stroke scale score were missing for 272 patients, leaving 29 228 for analysis adjusted for age and baseline severity (fig 1).

Statistical analysis
We compared outcome at 90 days in patients who received intravenous thrombolysis and controls for the whole cohort. We repeated the comparison among patients aged ≤80 and >80 years. We then examined the association of thrombolysis treatment with outcome within various age groups (<21, 21-30, 31-40, 41-50, 51-60, 61-70, 71-80, 81-90, and 91-100) to illustrate the strength of evidence across the full age range.

For each comparison, we looked at the overall distribution of all seven categories of scores on the modified Rankin scale in the two groups at day 90. The modified Rankin scale is an ordinal score used as a measure of functional outcomes in patients with stroke (see box).30 31

Scores on modified Rankin scale

0=no symptoms from stroke
1=no severe disability, despite symptoms
2=slight disability in which patients are unable to do all previous activities but able to look after themselves without help
3=moderate disability that requires some help, but patients can walk by themselves
4=moderately severe disability in which patients are unable to walk without assistance and need help for bodily needs
5=bedbound patients who are incompetent or require personal attention
6=death
To test for a significant association of distribution of outcome with exposure to thrombolysis we used the Cochran-Mantel-Haenszel statistic, adjusting for age and baseline score on the National Institutes of Health stroke scale as continuous variables.\textsuperscript{32,33} We had two reasons for our choice of baseline factors for adjustment. Firstly, age and baseline severity measured by National Institutes of Health stroke scale are the two most powerful prognostic factors for stroke outcome and are usually included in analyses of outcome distribution.\textsuperscript{32-36} Secondly, we had data on age and the stroke scale for our entire sample, whereas data on other factors of potential interest were incomplete. We also undertook a sensitivity analysis by considering the combined effect of the variables that differed significantly at baseline.

Our objective was mainly to undertake an ordinal distribution or “shift” analysis, which is an efficient endpoint analysis technique accepted by the European Agency for the Evaluation of Medicinal Products (EMEA).\textsuperscript{29} Shift analysis is considered to be better than dichotomisation of endpoint measures, though there are differences of opinion.\textsuperscript{38-43} Dichotomisation is criticised for the statistical information it discards, whereas shift analysis is especially useful when the treatment effect is mild or uniform, or both, across all Rankin categories, though in larger datasets it can sometimes incorrectly seem to violate proportionality assumptions.\textsuperscript{38-44} This is because the test for proportionality assumptions is described as “sensitive to sample size, such that large samples may produce statistically significant P values when in fact there is little practical difference between the cut-point-specific estimates.”\textsuperscript{45} Hence, though we planned to undertake examination of outcomes by proportional odds logistic regression analysis we also elected to undertake secondary analyses, dichotomising the Rankin scores 0-1 v rest, 0-2 v rest, and dead v survivors, to allow comparability of our findings with the other published data.\textsuperscript{46} Odds ratios in our analysis express the common odds of an improved distribution of outcome in association with treatment with alteplase.

The Cochran-Mantel-Haenszel test and proportional odds logistic regression analysis were undertaken with SAS 9.2 software and other analyses by StatsDirect software. We describe our findings in accordance with the STROBE guidelines.\textsuperscript{47}

Reliable information on symptomatic intracerebral haemorrhage was not available from VISTA controls as post-treatment imaging was not routinely carried out in neuroprotection trials in patients who had not been treated with alteplase. However, we compared the rates of symptomatic intracerebral haemorrhage for the definitions used within SITS (local or remote parenchymal haemorrhage type 2 on the imaging scan 22-36 hours after treatment, combined with a neurological deterioration of 4 or more points on the National Institutes of Health stroke severity scale from baseline, or from the lowest National Institutes of Health stroke severity score between baseline and 24 hours, or leading to death) and the NINDS study (any intracranial haemorrhage in the post-thrombolysis imaging scans if it was not seen on a previous imaging scan and any decline in neurological status) for younger and older patients of the SITS-ISTR registry.

**RESULTS**

All stroke patients were treated as per institutional practice and stroke guidelines acceptable at the point of their treatment. The table shows the baseline characteristics. Baseline severity of stroke was similar between patients who did and did not undergo thrombolysis among those aged \(\geq 80\) (P=0.6) and \(\leq 80\) (P=0.3).

Independently, baseline National Institutes of Health stroke scale accounted for 25.5% and age for 7.4% of the variation in 90 day outcome by modified Rankin scale (both P<0.001) and were included in all models, together explaining 29.6% of the variation.

**Overall outcome**

Across our whole sample, the distribution of scores on the modified Rankin scale at three months was better—that is, more patients had lower scores—among those who underwent thrombolysis (fig 2). The overall odds ratio was 1.6 (95% confidence interval 1.5 to 1.7; P<0.001) (fig 3).

**Outcomes among patients aged \(\leq 80\)**

Treatment with thrombolysis was associated with a significantly more favourable distribution of scores on the modified Rankin scale at three months (fig 2). The adjusted odds ratio was 1.6 (1.5 to 1.7); P<0.001; n= 25 789 (fig 3). P for test of proportionality assumption was <0.05. The unadjusted odds ratio was 1.5 (1.4 to 1.6) (n=26 028). Dichotomised outcomes were also significantly more favourable for thrombolysed patients than controls (1.9 (1.7 to 2.0) for score 0-2 v 3-6 on
### Baseline characteristics of patients in SITS*-VISTA

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>Mean (SD)</td>
<td>67.1 (12.6)</td>
<td>70.1 (12.2)</td>
</tr>
<tr>
<td></td>
<td>Median (range)</td>
<td>69 (10-98)</td>
<td>72 (21-101)</td>
</tr>
<tr>
<td>SBD</td>
<td>Mean (SD)</td>
<td>65.3 (11.6)</td>
<td>66.5 (10.7)</td>
</tr>
<tr>
<td></td>
<td>Median (range)</td>
<td>68 (10-80)</td>
<td>69 (21-80)</td>
</tr>
<tr>
<td>&gt;80</td>
<td>Mean (SD)</td>
<td>84.4 (3.25)</td>
<td>84.84 (3.38)</td>
</tr>
<tr>
<td></td>
<td>Median (range)</td>
<td>84 (81-98)</td>
<td>84 (81-101)</td>
</tr>
</tbody>
</table>

**No (%) of men**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>13,994/23,334</td>
<td>327/1,616</td>
<td>0.001</td>
</tr>
<tr>
<td>SBD</td>
<td>12,744/21,099</td>
<td>278/4,929</td>
<td>0.001</td>
</tr>
<tr>
<td>&gt;80</td>
<td>850/2,235</td>
<td>488/1,237</td>
<td>0.41</td>
</tr>
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</table>

**No (%) who had previously taken antithrombotics**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>8,776/22,792</td>
<td>1,267/2968</td>
<td>0.001</td>
</tr>
<tr>
<td>SBD</td>
<td>7,537/20,623</td>
<td>977/2414</td>
<td>0.001</td>
</tr>
<tr>
<td>&gt;80</td>
<td>1,239/2169</td>
<td>290/554</td>
<td>0.04</td>
</tr>
</tbody>
</table>

**No (%) with known diabetes mellitus**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>3,962/22,968</td>
<td>1,449/5,896</td>
<td>0.001</td>
</tr>
<tr>
<td>SBD</td>
<td>3,570/20,784</td>
<td>1,203/4,704</td>
<td>0.001</td>
</tr>
<tr>
<td>&gt;80</td>
<td>392/2,184</td>
<td>246/1,192</td>
<td>0.06</td>
</tr>
</tbody>
</table>

**No (%) with previous stroke**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>3,005/23,013</td>
<td>2,014/5,993</td>
<td>0.001</td>
</tr>
<tr>
<td>SBD</td>
<td>2,629/20,840</td>
<td>1,521/4,776</td>
<td>0.001</td>
</tr>
<tr>
<td>&gt;80</td>
<td>376/2,173</td>
<td>493/1,217</td>
<td>0.001</td>
</tr>
</tbody>
</table>

**No (%) with congestive heart failure**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>1,932/22,840</td>
<td>277/3,167</td>
<td>0.59</td>
</tr>
<tr>
<td>SBD</td>
<td>1,581/20,677</td>
<td>185/2,579</td>
<td>0.39</td>
</tr>
<tr>
<td>&gt;80</td>
<td>351/2,143</td>
<td>92/588</td>
<td>0.67</td>
</tr>
</tbody>
</table>

**No (%) with hypertension**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>14,331/22,875</td>
<td>4,170/5,896</td>
<td>0.001</td>
</tr>
<tr>
<td>SBD</td>
<td>12,687/20,683</td>
<td>3,273/4,704</td>
<td>0.001</td>
</tr>
<tr>
<td>&gt;80</td>
<td>1,644/2,192</td>
<td>897/1,192</td>
<td>0.87</td>
</tr>
</tbody>
</table>

**No (%) with atrial fibrillation**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Thrombolysis</th>
<th>Control</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>5,835/22,753</td>
<td>1,712/5,896</td>
<td>0.001</td>
</tr>
<tr>
<td>SBD</td>
<td>4,837/20,613</td>
<td>1,147/4,704</td>
<td>0.19</td>
</tr>
<tr>
<td>&gt;80</td>
<td>998/2,140</td>
<td>565/1,192</td>
<td>0.67</td>
</tr>
</tbody>
</table>

*Of patients with baseline National Institutes of Health stroke scale score in SITS datasheet, two (0.32%) in age group 31-40, five (0.30%) in age group 41-50, nine (0.25%) in age group 51-60, six patients (0.1%) in age group 61-70, eight (0.09%) in age group 71-80, and two (0.1%) in age group 81-90 were coded as having baseline score of 0 and treated with alteplase. They were assumed to have neurological deficit considered potentially disabling but not measured by restricted rules of the scale (such as distal limb weakness). These few cases will have no material impact on findings.

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Outcomes among patients aged >80

Among the 3439 patients aged over 80 with a 90 day score on the modified Rankin scale and baseline data on stroke severity, treatment with thrombolysis was associated with a significantly more favourable distribution of scores at three months compared with controls (fig 2). The adjusted odds ratio was 1.4 (1.3 to 1.6); P<0.001; n=3439 (fig 3). P for test of proportionality assumption was <0.05. The unadjusted odds ratio was 1.4 (1.2 to 1.6), (P<0.001; n=3472). Dichotomised outcomes were significantly more favourable for thrombolysed patients than controls (2.1 (1.7 to 2.5) for favourable outcome (score 0-2 on modified Rankin scale); 1.9 (1.5 to 2.3) for excellent outcome (score 0-1); and 0.89 (0.76 to 1.04) for mortality).

Our sensitivity analysis, in which we adjusted for age, sex, history of either diabetes or previous stroke, previous use of antithrombotics, baseline National Institutes of Health stroke severity score, and hypertension, yielded Cochran-Mantel-Haenszel P=0.003 and proportional odds of 1.5 (1.3 to 1.8) in favour of thrombolysis.

The results indicate that 8.2 patients aged >80 need to be treated for one more patient to achieve a modified Rankin scale score of 0-2.

Association of thrombolysis with outcome by age groups

Distributions of scores on the modified Rankin scale at 90 days were significantly better among thrombolysed patients than controls within each 10 year age group from 40 to 90 and, except among the small samples of patients younger than 30 and older than 90, point estimates for the adjusted odds ratios were consistent across all age groups (fig 3).

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**Modified Rankin scale**

<table>
<thead>
<tr>
<th>Score</th>
<th>Control (n=6160)</th>
<th>Alteplase (n=23334)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>21.8 (16.9)</td>
<td>12.7 (13.2)</td>
</tr>
<tr>
<td>1</td>
<td>12.8 (11.9)</td>
<td>10.8 (10.3)</td>
</tr>
<tr>
<td>2</td>
<td>11.9 (10.9)</td>
<td>9.3 (9.2)</td>
</tr>
<tr>
<td>3</td>
<td>15.8 (14.2)</td>
<td>8.3 (7.1)</td>
</tr>
<tr>
<td>4</td>
<td>13.9 (12.9)</td>
<td>8.7 (7.6)</td>
</tr>
<tr>
<td>5</td>
<td>10.9 (9.5)</td>
<td>13.6 (10.4)</td>
</tr>
<tr>
<td>Death</td>
<td>16.9 (17.1)</td>
<td>15.6 (16.6)</td>
</tr>
</tbody>
</table>

**Age >80**

<table>
<thead>
<tr>
<th>Score</th>
<th>Control (n=4929)</th>
<th>Alteplase (n=21099)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>12.1 (12.2)</td>
<td>12.8 (14.2)</td>
</tr>
<tr>
<td>1</td>
<td>11.1 (13.0)</td>
<td>12.2 (13.2)</td>
</tr>
<tr>
<td>2</td>
<td>12.2 (12.9)</td>
<td>9.5 (9.7)</td>
</tr>
<tr>
<td>3</td>
<td>10.7 (10.9)</td>
<td>32.6 (35.3)</td>
</tr>
<tr>
<td>4</td>
<td>11.3 (12.9)</td>
<td>35.3 (36.8)</td>
</tr>
<tr>
<td>5</td>
<td>15.1 (15.1)</td>
<td>44.8 (45.0)</td>
</tr>
</tbody>
</table>

**Fig 2** Scores on modified Rankin scale (from 0=no symptoms from stroke to 6=death) at three months between patients who underwent thrombolysis with alteplase and controls, indicating shift towards improved outcomes with thrombolysis. Numbers within coloured cells are percentages.
at 90 days
thrombolysis in VISTA neuroprotection trials; they achieved modified Rankin score of 0 and 4
score of 3, and two a score of 4; two died. Two patients aged 101 did not undergo
10 patients attained a score of 1, eight patients reached a score of 2, one patient achieved a
SITS and underwent thrombolysis; 15 patients reached a 90 day modified Rankin score of 0,
Controls
Favours
alteplase
Favours
control
Odds ratio
(95% CI)
0.2
0.5
1
2
5
<0.001
<0.001
1.9 (1.5 to 2.3)
1.7 (1.5 to 1.9)
1.7 (1.5 to 2.0)
1.5 (1.3 to 1.7)
1.3 (0.83 to 2.1)
1.3 (1.0 to 1.7)
1.5 (1.3 to 1.8)
1.5 (0.89 to 2.9)
1.3 (0.96 to 1.8; P=0.07). The corresponding rate for symptomatic intracerebral haemorrhage per National Institute of Neurological Disorders and Stroke definition27 (any increase in National Institutes of Health stroke scale from baseline and any parenchymal intracerebral haemorrhage) was significantly higher: 11.0% (229/20877) vs 8.3% (1670/20220); 1.4, 1.2 to 1.6; P<0.001).

Onset to treatment time
We calculated the time from onset of stroke to treatment for the administration of thrombolysis to patients in SITS-ISTR. The median time was similar in younger (≤80) and older (>80) patients (145 minutes, P=0.25 for
difference). Data on onset to treatment time for use of alteplase were not collected in VISTA.

DISCUSSION
Principal findings
Comparing patients from SITS who were treated with alteplase at an average of 145 minutes after stroke onset against controls from VISTA who received no alteplase we found more favourable outcomes with alteplase across the entire range of scores on the modified Rankin scale (odds ratio 1.6, 95% confidence interval 1.5 to 1.7, P<0.001). The nature and extent of this effect of alteplase is comparable with results from pooled analysis of randomised controlled trials, confirming the validity of our controlled but non-randomised analysis.4 We were able to examine outcomes separately among patients aged ≤80 and older patients aged >80. In each subgroup we found more favourable functional outcomes: odds ratios 1.6 (1.5 to 1.7), n=25 789, and 1.4 (1.3 to 1.6), n=3439, respectively.

Extending our analysis to smaller subgroups of age, we found independently significant benefits from alteplase in each 10 year age group from 40-90. We found no interaction between age and efficacy of alteplase and across the full age range from age under 20 to over 100. Only in patients aged under 30 did the trend not favour outcomes after use of alteplase.

In summary, we show that association between thrombolysis treatment and outcome is maintained in all patients, even in older patients, regardless of generally poorer outcomes in these age groups.

Strengths and limitations
Our analysis of SITS-VISTA data is based on almost 30 000 patients and confirms that there are improved outcomes after acute ischaemic stroke among patients who are offered thrombolytic therapy. The extent of the

Post-thrombolysis intracerebral haemorrhage
The rate of symptomatic intracerebral haemorrhage per SITS-MOST definition27 (≥4 point increase in National Institutes of Health stroke scale from baseline or death within 24 hours and parenchymatous haemorrhage (type PH2 or PHr2) at 22-36 hour imaging scans) was 2.5% (54/2163) among those aged >80 compared with 1.9% (398/20 759) among those aged ≤80, and thus not significantly higher (odds ratio 1.3, 0.96 to 1.8; P=0.07). The corresponding rate for symptomatic intracerebral haemorrhage per National Institute of Neurological Disorders and Stroke definition27 (any increase in National Institutes of Health stroke scale from baseline and any parenchymal intracerebral haemorrhage) was significantly higher: 11.0% (229/20877) vs 8.3% (1670/20220); 1.4, 1.2 to 1.6; P<0.001).

Fig 3 | Shift towards better outcomes on modified Rankin scale at three months adjusted for age and baseline severity (defined by National Institutes of Health stroke scale). Number of patients shown for age groups do not add up to 29 228 because numbers of patients ages <21 (n=38) and >100 (n=2) were too low to allow any comparison. All patients aged ≥21 were from SITS and underwent thrombolysis; 15 patients reached a 90 day modified Rankin score of 0, 10 patients attained a score of 1, eight patients reached a score of 2, one patient achieved a score of 3, and two a score of 4; two died. Two patients aged 101 did not undergo thrombolysis in VISTA neuroprotection trials; they achieved modified Rankine score of 0 and 4 at 90 days

Fig 4 | Odds ratios for score 0-1 on modified Rankin scale at three months adjusted for age and baseline National Institutes of Health stroke severity scale in patients who received thrombolytic therapy

<table>
<thead>
<tr>
<th>Age group</th>
<th>Odds ratio (95% CI)</th>
<th>Odds ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤80</td>
<td>1.6 (1.5 to 1.7)</td>
<td>2.2 (0.2 to 6.0)</td>
</tr>
<tr>
<td>&gt;80</td>
<td>1.9 (1.5 to 2.3)</td>
<td>1.6 (1.5 to 1.7)</td>
</tr>
<tr>
<td>All age groups</td>
<td>1.5 (1.5 to 1.7)</td>
<td>1.3 (0.36 to 4.6)</td>
</tr>
</tbody>
</table>

Odds ratios for score 0-1 on modified Rankin scale at three months adjusted for age and baseline National Institutes of Health stroke severity scale in patients who received thrombolytic therapy.
apparent benefit matches that from published randomised trials. These observations extend to older age groups and only in a small group of patients aged 91-100 (137 patients in alteplase group and 77 in the control group) did we fail to show significance, with consistent point estimates but wide confidence intervals. The point estimates for improved outcomes in this age group are also consistent with the published data.

We undertook our primary analyses using “shift analysis,” an analytical approach accepted by the European Drug Licensing agency. The Cochran-Mantel-Haenszel test is a non-parametric approach that avoids invoking an assumption of a common odds ratio (that is, proportionality) across all cut points on the ordinal outcome scale. It provides a conservative estimate of significance. Because it does not express the extent of the association, we also applied an ordinal logistic regression analysis to estimate a common odds ratio across categories of the modified Rankin scale. Again, we found significantly better outcomes, though the proportionality assumption was not satisfied. Whereas a non-significant result for proportionality would imply that common odds could be assumed, the converse does not necessarily apply.

The proportionality assumption test might be oversensitive when applied to large sample sizes. Furthermore, it is a global test that cannot differentiate the heterogeneity resulting from alteplase or other covariates. As our sample sizes were large we could still be justified in using the odds estimated from ordinal logistic regression. Even so, for final confirmation we used a less powerful dichotomised approach.

There were improved outcomes among patients who underwent thrombolyis in age groups from 31 to 90. We found no improved outcomes for patients aged under 30 and above 90, but the small number of patients in these groups greatly reduced statistical power for these analyses and the trends mostly followed the same pattern as for intermediate ages.

We chose age and baseline National Institutes of Health stroke scale score for adjusted analysis mainly because of their established roles of influence on stroke outcomes. We also undertook sensitivity analysis, adjusting for differences in age, sex, history of diabetes or previous stroke, previous use of an antithrombotic, baseline National Institutes of Health stroke scale score, and hypertension, between those who did and did not undergo thrombolysis. The adjusted analyses for these variables confirmed significant findings for improved outcomes with thrombolysis regardless of age.

The baseline demographic characteristics for the complete dataset favoured the thrombolysis group. This influence, however, did not extend to patients aged >80. As a result, though our estimates of overall effect of alteplase could be biased, the relative differences between subgroups should remain reliable.

Our conclusions derive merit from being based on a huge population of patients who were treated in routine clinical practice and compared against controls from rigorously conducted neuroprotection trials: any bias in quality of care should favour the control group. The limitation of SITS-ISTR data has been discussed extensively in previous publications. In short, SITS-ISTR is a registry, and it is therefore impossible to guarantee completeness of inclusions and to exclude selection bias. For a sample of patients included in SITS-MOST (monitoring study), source data were verified onsite by monitors under the

<table>
<thead>
<tr>
<th>Age group</th>
<th>Odds ratio (95% CI)</th>
<th>Odds ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>21-30</td>
<td>0.69 (0.08 to 5.8)</td>
<td>1.9 (1.16 to 3.1)</td>
</tr>
<tr>
<td>31-40</td>
<td>1.6 (1.2 to 2.1)</td>
<td>1.9 (1.6 to 2.2)</td>
</tr>
<tr>
<td>41-50</td>
<td>1.8 (1.6 to 2.0)</td>
<td>2.0 (1.8 to 2.2)</td>
</tr>
<tr>
<td>51-60</td>
<td>2.1 (1.7 to 2.5)</td>
<td>1.9 (0.81 to 4.4)</td>
</tr>
<tr>
<td>61-70</td>
<td>1.9 (1.7 to 2.0)</td>
<td>2.1 (1.7 to 2.5)</td>
</tr>
<tr>
<td>71-80</td>
<td>1.9 (1.7 to 2.0)</td>
<td>2.1 (1.7 to 2.5)</td>
</tr>
<tr>
<td>81-90</td>
<td>1.9 (1.8 to 2.1)</td>
<td></td>
</tr>
<tr>
<td>91-100</td>
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</table>

All age groups 1.9 (1.8 to 2.1)
WHAT IS ALREADY KNOWN ABOUT THIS TOPIC

In the European Union patients with ischaemic stroke aged >80 are not recommended for thrombolysis, though no one has shown negative outcomes. Age is a non-modifiable risk factor for stroke, and the proportion of older patients is rapidly rising in our society.

WHAT THIS STUDY ADDS

Odds for improved outcomes in patients with ischaemic stroke who receive thrombolysis (with alteplase) are similar in those aged ≤80 and >80. Age alone should not be a barrier to thrombolysis.

supervision of the relevant national coordinator. The monitors could examine admissions department records, etc, to check for completeness of data. Individual investigators’ results are not published, limiting the incentive for selection; in contrast, the sharing of total enrolment numbers might act as an incentive to be inclusive. The almost identical main outcomes in SITS-MOST and randomised controlled trials after adjustment for baseline differences suggests that the influence from such potential bias is limited. Subsequent studies based on SITS-ISTR data also show the similar outcome for the overall study population compared with the SITS-MOST. Selective reporting of good outcomes in SITS could generate an opposite bias but was limited through site monitoring procedures used in the SITS-MOST study.

None of the neuroprotective agents used for the patients in the VISTA control group has an influence on outcome, and over half of the VISTA cohort received only placebo.

Because VISTA lacks data on repeat brain imaging among patients who did not receive thrombolysis, we had no data on symptomatic intracerebral haemorrhage in our control group. Therefore, we compared the rates between patients aged >80 and ≤80 only with SITS data. There was no difference in rates between those aged >80 and ≤80 with the SITS-MOST definition but slightly higher with the NINDS definition. In a complementary per protocol analysis of SITS-ISTR data (that is, patients’ selection based on SITS-MOST criteria except for its age criterion), there was no significant difference in rates of symptomatic intracerebral haemorrhage among patients aged >80 compared with the younger cohort (1.8% v 1.7%, P=0.70, adjusted odds ratio 0.90, 0.73 to 1.09). Our SITS patients in the SITS-VISTA dataset are unselected and therefore rates of symptomatic intracerebral haemorrhage are slightly higher in the current study.

Regardless, we have shown that even if there were any more haemorrhages among elderly patients who receive thrombolysis, based on a conservative definition, there seems to be no adverse influence on the distribution of outcomes. In fact, we observe a beneficial effect on mortality. Others have concluded that factors such as comorbidity, rather than use of alteplase, are responsible for the observed increase in late case fatality among older patients.21 34

Comparison with other studies

We reached the same conclusions as analyses that used VISTA data or the limited pooled randomised trial data in elderly patients. Elderly patients treated with thrombolysis in trials reported by VISTA (n=5817) had significantly better adjusted outcomes than patients who did not receive thrombolysis (odds ratio 1.3, 1.1 to 1.7; P=0.002). Elderly patients treated in the pooled randomised trials showed a trend towards better adjusted outcomes (score on modified Rankin scale 0-2 v 3-6) than those who did nor receive thrombolysis (1.8, 0.73 to 4.3; n=137). Previous studies have shown findings consistent with our results but on small datasets.

Despite these points, treatment allocation in our study was not randomised. More extensive data from randomised controlled trials could more conclusively answer this question. Two trials are currently examining this topic. A trial in Italy has so far enrolled around 17% of the planned 600 patients over a two year period. In the UK, the International Stroke Trial-3 (IST-3) is examining outcomes among all patients who receive thrombolysis and has no upper age limit in its inclusion criteria.

Over 12 years, the trial has enrolled around 2400 patients of the originally planned 6000. About a third of these patients are aged >80 and are being treated within the time window of interest.

Conclusions and policy implication

In our analysis, patients who were treated with intravenous alteplase had better outcomes than untreated patients, and this effect was not dependent on age. In particular, patients aged over 80 derived similar benefit from treatment as younger patients. The weight of evidence to date indicates a potential for benefit in the older people, and there is no a priori reason to suspect a diminished effect compared with younger people. Furthermore there are reassuring safety data on the risk of intracerebral haemorrhage. We conclude that clinical treatment guidelines should be revised to remove the age restriction in use of intravenous alteplase for acute ischaemic stroke. Age alone should not be a barrier to treatment.
authors take full responsibility for the content. The manuscript was reviewed and approved by the steering committees of VISTA and SITS. No commercial organisation was involved in the origination, execution or reporting of this work. NIKM is guarantor.

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Competing interests: All authors have completed the Unified Competing Interest form at www.icmje.org/col1_disclosure.pdf (available on request from the corresponding author) and declare: KRL has received honorariums from Boehringer Ingelheim, Lundbeck, Thrombogenics, Talecs. GA is a member of the advisory board for Boehringer Ingelheim, Denmark. JAE was investigator of ECASS II, ECASS III, and PROFESS promoted by Boehringer Ingelheim. NA is an employee of Siemens International, which received a grant from Boehringer Ingelheim for the SITS-MOST/SITS-ISTR study with alteplase. NGW has received expenses from Boehringer Ingelheim for his role as member of the steering committee in relation to the ECASS III trial with alteplase and served as a consultant to Thrombogenics as chairman of the DSMB. SITS International (chaired by NGW) received a grant from Boehringer Ingelheim and from Ferrer for the SITS-MOST/SITS-ISTR. His institution has also received grant support towards administrative expenses for coordination of the ECASS III trial. NGW has also received lecture fees from Boehringer Ingelheim and from Ferrer. PAR is the German deputy national SITS coordinator. He was investigator of ECASS III and PROFESS sponsored by Boehringer Ingelheim and of DIAS and DIAS-2 sponsored by PAION. He received honorariums and travel expenses from Boehringer Ingelheim, PAION, and Ferrer.

Ethical approval: Not required.

Data sharing: Data sharing: no additional data available.

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