The Albumin in Acute Stroke (ALIAS) Multicenter Clinical Trial
Safety Analysis of Part 1 and Rationale and Design of Part 2
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Background and Purpose—Enrollment in the Albumin in Acute Stroke (ALIAS) Trial was suspended in late 2007 due to a safety concern. We present the safety data of that Trial (“Part 1”) and the rationale for the design of Part 2.

Methods—ALIAS Part 1 was designed to assess whether 25% albumin (ALB) started within 5 hours of stroke onset would confer neuroprotection in subjects with acute ischemic stroke and baseline National Institutes of Health Stroke Scale of ≥6. Exclusion criteria included recent or current congestive heart failure, myocardial infarction, or cardiac surgery. The study comprised 2 cohorts: subjects who received thrombolysis and those who did not, each with 1:1 randomization to ALB or placebo. The primary outcome was the National Institutes of Health Stroke Scale and modified Rankin Scales at 90 days. The intended sample size was 1800.

Results—Four hundred thirty-four subjects were enrolled, and 424 were used in the safety analysis (ALB 207, saline 217). There were 36 deaths within the first 30 days in the ALB group and 21 in the saline group. In contrast, death rates after 30 days were similar by treatment. Large strokes were the predominant cause of early death in both groups. In subjects >83 years of age, 90-day death rates were 2.3-fold higher with ALB than with saline (95% CI, 1.04 to 5.12). Similarly, 90-day deaths in subjects receiving excessive fluids were 2.10-fold greater with ALB than with saline (CI, 1.10 to 3.98).

Conclusions—The ALIAS Part 2 Trial, which started in early 2009, was modified as follows to enhance safety: upper age limit of 83 years; requirement for normal baseline serum troponin level; restriction of total intravenous fluids in the first 48 hours to ≤4200 mL; mandatory diuretic at 12 to 24 hours; and detailed site retraining. Because of insufficient nonthrombolysed subjects (22%) in Part 1, the 2-cohort design was eliminated. The Data Safety Monitoring Board has reviewed the safety data of Part 2 3 times and has approved continuation of the trial. (Stroke. 2011;42:119-127.)

Key Words: albumin ischemic stroke neuroprotectant randomized controlled trial

In the clinical management of acute ischemic stroke, there is a compelling, unmet need for safe and effective neuroprotective strategies to limit brain injury, facilitate brain repair, and improve functional outcome.1 Extensive animal studies have shown human albumin (ALB) in moderate to high doses to be a promising neuroprotectant in focal and global cerebral ischemia and traumatic brain injury.2-7 In focal ischemia, ALB diminished total infarct volume by two thirds and reduced brain edema by three fourths or more with a therapeutic window of efficacy extending to 4 hours; ameliorated brain swelling2,3,6; improved blood flow to critically perfused brain regions6; enhanced microvascular perfusion6,10; reduced postischemic microvascular blood element adhesion11; and helped to transport important free fatty acids to the postischemic brain.12 A National Institute of Neurological Disorders and Stroke (NINDS)-funded Phase I pilot clinical trial was subsequently conducted in 82 acute ischemic stroke subjects who received 25% human albumin in doses that were escalated into the experimentally neuroprotective range.13,14 ALB therapy was safely tolerated: mild to moderate pulmonary edema occurred in 13% of subjects but responded readily to medical management. Exploratory efficacy analysis suggested that a beneficial treatment effect might exist.14 After the pilot trial, the Albumin in Acute Stroke (ALIAS) Trial (ClinicalTrials.gov Identifier NCT00235495) was begun as a randomized, double-blind, placebo-controlled trial whose primary aim was to ascertain whether high-dose ALB...
therapy (2 g/kg) administered within 5 hours of stroke onset would increase the proportion of subjects with favorable outcome at 3 months compared with saline-placebo.\textsuperscript{13} The ALIAS Trial was sponsored by the NINDS and operated under a Food and Drug Administration Investigational New Drug license. The first ALIAS subject was recruited in July 2006. In December 2007, at the first interim analysis of 3-month follow-up data in 225 subjects (but after 434 subjects had been enrolled) at 62 North American clinical sites, the trial’s independent Data and Safety Monitoring Board (DSMB) recommended to NINDS that enrollment be suspended due to safety concerns and that the study team consider revising the protocol so as to enable the trial to resume with increased safety. The ALIAS principals were granted permission to review the safety data in an unblinded manner. After extensive internal conferences and discussions with external advisors, the ALIAS Executive Committee developed a revised protocol and analysis plan together with a comprehensive site-training program. These changes were approved by the DSMB in July 2008 and by the Food and Drug Administration in September 2008. The ALIAS Trial then began as a separate, new study referred to as “Part 2” and enrolled its first subject in February 2009.

Revisions to the protocol were based on our unblinded analysis of the Part 1 safety data and were implemented in an effort to improve the safety profile of trial participants. This article presents the results of our evaluation of the safety data from Part 1 and sets forth the rationale for the design changes instituted in the currently ongoing Part 2.

**Methods**

The ALIAS Part 1 Trial was originally designed as 2 separate but concurrently implemented double-blind, Phase III multicenter trials with 1:1 randomization to ALB or saline-placebo. The objective was to assess whether 25% ALB therapy (2 g/kg intravenously administered over 120 minutes) compared with an equal volume of 0.9% normal saline conferred neuroprotection in acute ischemic stroke, over and above the standard of care, in 2 cohorts of patients with acute ischemic stroke. One cohort consisted of subjects who received standard-of-care thrombolytic therapy (intravenous tissue plasminogen activator [tPA], intra-arterial tPA, endovascular mechanical thrombolyis with approved devices and catheters, or a combination of intravenous and endovascular treatment). The other cohort consisted of subjects who were not thrombolysed. The administration of thrombolytic therapy was based on local clinical judgment informed by then-current guidelines.\textsuperscript{16} The rationale for this design stemmed in part from preclinical evidence that ALB did not require induced reperfusion to confer neuroprotection\textsuperscript{17} and from an observed trend in the ALIAS Pilot Trial of better 3-month outcome with higher doses of ALB in both the tPA and the non-tPA cohorts.\textsuperscript{18} The study design of the ALIAS Part 1 Trial was identical for the 2 cohorts (thrombolysis and nonthrombolysis). The eligibility criteria are presented in Table 1.

A centralized step-forward, web-based 1:1 randomization process was used. All study personnel and patients were blinded. A biased-coin minimization algorithm adjusted for clinical site within each cohort.\textsuperscript{19} Study-drug kits consisted of a 500-mL and a 250-mL bottle of either 25% ALB or saline encased in blinding boxes and delivered through tinted intravenous tubing.\textsuperscript{18} Albumin was manufactured for the trial by Baxter Healthcare Corp, Westlake Village, Calif. A bedside nurse or other personnel not involved with the trial administered the study drug (8 mL/kg) by constant intravenous infusion over 2 hours (±15 minutes). Subjects weighing ≥94 kg received a maximum volume of 750 mL.

**Table 1. Inclusion and Exclusion Criteria, ALIAS Part 1 Trial**

<table>
<thead>
<tr>
<th>Inclusions</th>
<th>Exclusions</th>
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<tbody>
<tr>
<td>Acute ischemic stroke</td>
<td>Any findings of CHF on physical examination (jugular venous distention, third heart sound, resting tachycardia &gt;100 beats/min attributable to CHF, abnormal hepatojugular reflux, lower extremity pitting edema attributable to CHF or unexplained, bilateral rales); and/or definite evidence of pulmonary edema on chest x-ray (if performed; not required)</td>
</tr>
<tr>
<td>Age ≥18 years</td>
<td>Current acute or chronic lung disease requiring supplemental O₂ therapy</td>
</tr>
<tr>
<td>Baseline NIHSS score of ≥6 as assessed immediately before tPA treatment in the thrombolytic cohort or immediately before randomization in the nonthrombolytic cohort</td>
<td>History of or known allergy to ALB or to natural rubber latex</td>
</tr>
<tr>
<td>Initiation of ALB/placebo treatment within 5 hours of stroke onset and within 60 minutes of the start of intravenous tPA if given</td>
<td>Pregnancy (women of childbearing age must have a negative pregnancy test)</td>
</tr>
<tr>
<td>Signed and dated informed consent</td>
<td>Concurrent participation in any other therapeutic clinical trial</td>
</tr>
<tr>
<td>Suspected or symptoms of acute MI on admission</td>
<td>Evidence of any other major life-threatening or serious medical condition that would impair completion of the trial, impair outcome assessment, or in which ALB therapy might be contraindicated or harmful</td>
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</table>

Vital signs were monitored frequently during and after study-drug administration. Serum chemistries were collected at 24 and 48 hours. Intravenous fluid intake was recorded at 24 and 48 hours. A follow-up brain CT or MRI scan was obtained at 24 hours. An electrocardiogram was repeated at 24 to 48 hours. Neurological and cardiac status, including National Institutes of Health Stroke Scale (NIHSS) score, was assessed at 24 and 48 hours and at 7 days or discharge, whichever came first. Diuretic treatment was not mandated, but administration of a loop diuretic such as furosemide in an initial dose of 10 to 20 mg intravenously was recommended if clinically indicated. Antiplatelet therapy was recommended in all subjects within 48 hours of their stroke. Blood pressure was managed according to the local standard of care.

Subjects were followed for 1 year. At 3 months (±14 days) postrandomization, subjects were required to come to the clinic, where the NIHSS, modified Rankin Scale score, Barthel Index, Stroke-Specific Quality of Life instrument,\textsuperscript{19} and Trailmaking A and B\textsuperscript{19} were assessed by a site investigator who was certified in
Table 2. Baseline Characteristics of the Safety Cohort

<table>
<thead>
<tr>
<th></th>
<th>Albumin (N=207)</th>
<th>Saline (N=217)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>69.3±13.6 (SD) years (maximum, 97)</td>
<td>69.8±14.6 (SD) years (maximum, 97)</td>
</tr>
<tr>
<td>Gender</td>
<td>56.0% male</td>
<td>50.2% male</td>
</tr>
<tr>
<td>Race</td>
<td>86.5% white, 7.7% black</td>
<td>81.6% white, 12.0% black</td>
</tr>
<tr>
<td>Time from stroke onset to study-drug treatment</td>
<td>202±50 minutes</td>
<td>206±50 minutes</td>
</tr>
<tr>
<td>Time from stroke onset to intravenous tPA</td>
<td>137±33 minutes</td>
<td>138±36 minutes</td>
</tr>
<tr>
<td>Baseline NIHSS score</td>
<td>Median, 11. NIHSS 6–10, 43.5%; 11–15, 23.7%; 16–20, 17.9%; 21–25, 9.7%; &gt;25, 5.3%</td>
<td>Median, 11. NIHSS 6–10, 42.4%; 11–15, 24.0%; 16–20, 18.9%; 21–25, 11.1%; &gt;25, 3.7%</td>
</tr>
<tr>
<td>Medical history</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>73.4%</td>
<td>75.1%</td>
</tr>
<tr>
<td>Ischemic heart disease</td>
<td>15.9%</td>
<td>18.4%</td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>21.7%</td>
<td>24.9%</td>
</tr>
<tr>
<td>Previous stroke; previous transient ischemic attack</td>
<td>16.9%; 10.6%</td>
<td>18.4%; 15.2%</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>20.3%</td>
<td>22.6%</td>
</tr>
<tr>
<td>Oxfordshire stroke classification</td>
<td>50.2%</td>
<td>54.0%</td>
</tr>
<tr>
<td>Partial anterior circulation</td>
<td>50.2%</td>
<td>54.0%</td>
</tr>
<tr>
<td>Total anterior circulation</td>
<td>30.0%</td>
<td>27.4%</td>
</tr>
<tr>
<td>Lacunar</td>
<td>13.5%</td>
<td>11.6%</td>
</tr>
<tr>
<td>Posterior circulation</td>
<td>5.8%</td>
<td>6.5%</td>
</tr>
<tr>
<td>Baseline systolic blood pressure</td>
<td>160±29 mm Hg</td>
<td>157±29 mm Hg</td>
</tr>
<tr>
<td>Baseline plasma glucose</td>
<td>7.4±3.3 mmol/L</td>
<td>7.5±2.9 mmol/L</td>
</tr>
<tr>
<td>CT-based ASPECTS score of 0 to 7 (central reader)</td>
<td>33.2%</td>
<td>27.7%</td>
</tr>
</tbody>
</table>

ASPECTS indicates Alberta Stroke Programme Early CT Score.

outcomescale completion and blinded to the subject’s admission treatment assignment and hospital course. Subjects were also followed by telephone contact at 1 month (±7 days), 6 months (±14 days), 9 months (±14 days), and 1 year (±14 days) postrandomization to assess the modified Rankin Scale, record serious adverse events (SAEs), and complete the EuroQol22 at 3 months and 1 year and the Questionnaire to Validate a Stroke-Free Status22 at 3, 6, 9, and 12 months. We assessed blinding by asking the rater to indicate what treatment assignment they thought the patient had received. The raters’ responses were correct 52.4% of the time (177 correct out of 338 responses), indicating that the outcome assessment had been truly blinded.

A favorable outcome was defined as an NIHSS score of 0 to 1 and/or a modified Rankin Scale score of 0 to 1 at 90 days postrandomization. With a 2-sided Type I error probability of 0.05, power of 80% to detect a 10% absolute effect-size difference in the primary outcome, and an assumption of the control group’s favorable outcome proportion to be 40%, the required sample size was 900 in each cohort, or a total of 1800 subjects. Because the primary analysis was based on the intent-to-treat principle, the sample size included inflation to account for crossovers and missing data as well as for 3 interim analyses for overwhelming efficacy or futility.

Due to premature suspension of the trial after 434 subjects had been enrolled, neither the thrombolytic (N=327) nor the nonthrombolytic cohort (N=97) had sufficient power to test the primary hypothesis. Thus, we evaluated the safety data by combining the cohorts.

Results

There were 434 randomized subjects, 215 allocated to ALB and 219 to saline treatment. Of these, 424 received at least 20% of the study drug (ALB 207, saline 217) and were used in the safety analysis. The baseline characteristics of the safety cohort, shown in Table 2, were very similar to those of the entire population. (A CONSORT diagram of the Part 1 Trial is available online as a Supplementary Figure; available at http://stroke.ahajournals.org.)

The Figure presents the major safety events by treatment assignment. As anticipated, pulmonary edema occurred approximately 3-fold more often in ALB- than in saline-treated subjects; the event rate of 12.1% in ALB subjects is similar to the 13% incidence observed in the ALIAS Pilot Trial.13 In the thrombolytic cohort, the proportion of subjects with symptomatic intracranial hemorrhage was similar in ALB (4.2%) and saline subjects (5.2%) and similar to that of the NINDS tPA trial.18 Symptomatic hemorrhage was defined as the occurrence of intracranial hemorrhage within 24±6 hours of randomization, proven by neuroimaging (MRI or CT), and associated with deterioration in neurological status. In the investigator’s opinion, the hemorrhage must have been thought to be the primary cause of the subject’s deterioration. Although the DSMB did not disclose the details of its confidential deliberations, we believe that its recommendation to suspend subject recruitment in ALIAS Part 1 was based primarily on an observed imbalance in overall deaths in the 2 groups (Figure; Table 3). Deaths in the ALB and saline groups were similar on Days 1 to 4 after randomization,
whereas between Days 5 and 30 there were more deaths in the ALB group compared with the saline group (Table 3). By contrast, death rates beyond 30 days were virtually identical in the 2 treatment groups. Safety data of all subjects who died were reviewed in a treatment-blinded fashion (by M.D.G. and M.D.H.) to assign a primary cause of death. Large strokes (with or without medical complications) were the predominant cause of death throughout Days 1 to 30, and these were more frequent in ALB- than in saline-treated subjects (Table 3). No single cause of death, however, completely explained the difference in deaths by treatment assignment.

To assess potential treatment-related factors, we conducted univariate analyses of 90-day deaths for various baseline variables. The relative risks (RRs) and 95% CIs are: age (RR, 1.04; CI, 1.02 to 1.06), baseline NIHSS (1.13; 1.09 to 1.17), plasma glucose (1.03; 0.97 to 1.10), baseline Alberta Stroke Programme Early CT Score of 8 to 10 versus 0 to 7 (0.34; 0.21 to 0.55), onset to study-drug treatment (1.00; 0.99 to 1.01), and cohort (1.25; 0.67 to 2.33). A multivariable model incorporating treatment, baseline NIHSS, and age showed a nonsignificant effect of treatment (1.49; 0.93 to 2.38) but significant effects of baseline NIHSS (1.10; 1.06 to 1.14) and age (1.03; 1.01 to 1.05).

### Influence of Advanced Age and Fluid Excess

Because the multivariable analysis showed a significant treatment effect for age, we explored the differential death rate in the safety cohort by various dichotomized age groups beginning at age 80 years. Ninety-day death rates did not differ significantly in ALB and saline subjects aged ≤83 years, whereas 90-day deaths in subjects >83 years were 2.3-fold higher in the ALB than in the saline group (Table 4A). We also hypothesized that differences in intravenous fluids may have contributed to ALB-associated deaths. We considered fluid excess to be present if a subject received >4200 mL of total intravenous fluids during the first 48 hours (based on an assumed 75 mL/hr intravenous fluid rate plus the volumes of study drug and tPA). Among subjects without excessive fluids, 90-day death rates were not significantly associated with treatment assignment, whereas among subjects with fluid excess, death rates were 2-fold greater in ALB than in saline subjects (RR, 2.10; CI, 1.10 to 3.98; Table 4A).

### Table 3. Deaths by Treatment Assignment, Day, and Adjudicated Cause of Death

<table>
<thead>
<tr>
<th></th>
<th>Days 1– 4</th>
<th></th>
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<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>ALB</td>
<td>Saline</td>
<td>ALB</td>
<td>Saline</td>
<td>ALB</td>
<td>Saline</td>
<td>ALB</td>
</tr>
<tr>
<td>Large stroke</td>
<td>6</td>
<td>4</td>
<td>8</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Large stroke + complications</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>ICH as primary cause</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary cardiac cause</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
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<td></td>
<td></td>
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<tr>
<td>Medical complication</td>
<td>1</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other known cause</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unknown cause</td>
<td>1</td>
<td></td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Column totals</td>
<td>9</td>
<td>9</td>
<td>15</td>
<td>6</td>
<td>8</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

ICH indicates intracranial hemorrhage.
4B). However, because fluid administration occurred during and after study-drug administration, it is possible that excess fluid volume is a merely a marker of worse outcome rather than its cause.

These results suggested the potential for minimizing excess deaths by attention to these 2 factors. This was modeled by comparing the 90-day death rates in the entire safety cohort versus the subgroup with out-of-hospital stroke, age >84, and with 48-hour intravenous fluids >4200 mL (Table 4C). Subjects with in-hospital strokes were excluded based on our impression that these patients tended to be more ill at baseline and to have more adverse events. The result suggested that implementation of the age and fluid restrictions and the exclusion of in-hospital strokes would eliminate significant treatment-related differences in 90-day deaths.

### Serious Adverse Events

SAEs occurred in 53.6% of ALB subjects and 46.5% of placebo subjects. Cardiovascular SAEs (CV-SAEs) were coded more commonly in ALB subjects (21.7%) than in the placebo group (8.3%; RR, 2.60; 95% CI, 1.57 to 4.37), primarily due to higher SAE rates of pulmonary edema (6.8% versus 2.8%) and acute coronary syndrome (8.2% versus 0.5%) in ALB-treated subjects than in saline-treated subjects.

Among subjects with CV-SAEs, myocardial infarction (MI) was diagnosed in 15 ALB-treated subjects (33%) but in only 2 saline-treated cases (11%). Of the 15 ALB-treated subjects with MI, 7 were diagnosed acutely on Days 1 or 2; 6 of those subjects died, although in only 2 of these cases could the death be directly attributed to an acute cardiac cause (ie, progressive hypoxemia, hemodynamic instability leading to shock). Among those CV-SAE subjects without a clinical diagnosis of MI on Days 1 to 2, elevated serum troponin levels at 24 and/or 48 hours were noted in 11 of 39 ALB subjects (28%) but in only 1 of 16 saline-treated subjects (6%). Although we concluded that cardiopulmonary pathophysiology must have played a role in the excess mortality, we were unable to show convincingly that it was the sole or specific cause.

### Redesign of ALIAS Trial, Part 2

Part 2 of the ALIAS Trial, a standalone study, retains many structural design features of the Part 1 Trial; that is, it is a
concurrently controlled, parallel 2-arm trial of ALB versus saline with a 1:1 randomization ratio. The primary efficacy measure remains unchanged, and we maintain 40% as the control group’s presumed proportion of good outcome. The major modifications from the Part 1 protocol are listed subsequently and the rationale for their implementation discussed.

- Upper age limit of 83 years at the time of randomization: See Table 4a.
- Requirement that baseline serum troponin level be ≤0.1 μg/L. Elevated baseline troponin levels may occur in a minority of ischemic stroke, denote some degree of cardiac injury, and may therefore be associated with cardiac adverse events. The ALIAS DSMB recommended that troponin levels be reviewed before randomizing a patient in the ALIAS Part 2 Trial.
- Exclusion of patients with in-hospital strokes. These patients have significant comorbid illness and are more likely to suffer SAES and less likely to respond to treatment.
- Imposition of strict intravenous fluid management guidelines: total intravenous fluids in first 48 hours not to exceed 4200 mL (unless cogent medical indications exist); strict monitoring of intravenous fluid intake; and the mandatory administration of a loop diuretic (typically furosemide, 20 mg intravenously) between 12 and 24 hours after study-drug treatment. Age is associated with loss of left ventricular compliance with associated diastolic dysfunction and, therefore, potentially reduced ability to compensate in the face of a fluid challenge, particularly a prolonged one such as is seen with high-dose ALB administration.
- Implementation of a detailed retraining module for clinical site staff, including a mandatory certification test and site principal investigator attestation form. Mandatory retraining (for all clinical sites that participated in Part 1) and mandatory training (for clinical sites that did not participate in Part 1) was required for participation in Part 2. A web-based training and testing module was developed. Each clinical site principal investigator provided a signed attestation that he or she would provide retraining to all involved clinical personnel.
- Inclusion of baseline NIHSS score as a covariate in the primary efficacy and safety analyses. The clinical trials literature has repeatedly cited the benefits of covariate-adjusted analysis for improving statistical power, particularly for covariates that are highly correlated with the outcome measure.24,25
- Combining subjects with and without thrombolytic treatment into a single cohort: inclusion of thrombolysis treatment status in the primary efficacy analysis model. The original premise of the ALIAS Trial, that ALB is effective in patients who received thrombolytic treatment as well as those who did not, has not changed. However, we assumed that we would see a greater effect in the former group (possibly due to a synergistic effect of tPA and ALB14) and, hence, we had designed Part 1 of the trial to conduct separate studies in these 2 cohorts of patients. Because of the insufficient enrollment into Part 1 of subjects who did not receive thrombolytic treatment, we are combining the 2 groups in Part 2. It is possible that we may observe a modest effect (eg, 5% to 10% treatment effect) in the nonthrombolysis stratum and a very large effect (eg, 20% to 30%) in the thrombolysis stratum. In such a scenario, we might find a statistically significant interaction effect; however, we would wish to conclude that there is a significant study treatment effect overall. Hence, we shall consider a statistically significant interaction effect only if it is also qualitative, that is, if the treatment effect is in the opposite direction in the 2 strata. In such a case, the primary efficacy analysis will be based on the thrombolysis stratum only.

We conducted multiple simulations to determine the sample size needed to address adequately the power for the interaction effect and the main study treatment effect for the entire study as well as for the thrombolysis stratum only. We concluded that a total sample size of 1100 will provide sufficient power (80%) and minimize Type I error probability for the overall Part 2 Trial. The clinically significant interaction effect is defined as a 20% differential treatment effect between the thrombolysis and nonthrombolysis strata. We believe that this value of 20% is justified because in the ALIAS Pilot Trial,14 a 26.1% absolute difference in good outcome occurred in the high-dose ALB tiers of the thrombolysis and nonthrombolysis cohorts. The sample size of 1100 was determined via simulation to ensure that the 20% interaction effect could be detected with 80% power at a 2-sided α=0.10.
- Inclusion of Statistical Safety Monitoring Guideline. In addition to stopping guidelines based on efficacy and futility, we established a statistical stopping guideline for safety based on the 30-day mortality rate. The rationale for basing the stopping guidelines on the number of events (referred to as reverse sampling method) rather than on the number of subjects is because we are unsure of the precise estimate of the event rates. Safety assessments based on the number of subjects may yield a decision-making process based on unstable estimates, because a relatively small (approximately 10%) event rate is anticipated in the control group. With 100 subjects enrolled in the study, for instance, a treatment group differential of only 1 death (which can happen by chance) would exaggerate the relative risk unnecessarily.
- Planned Meta-Analysis of Parts 1 and 2. After completion of Part 2 and analysis of its data, we plan to conduct a meta-analysis of Parts 1 and 2 using summary statistics from the 2 cohorts of Part 1 and the Part 2 study cohort weighted using the inverse normal method. We also plan to conduct pooled analysis of individual data from both parts adjusting for the study and cohort/strata effect.

**Current Status of ALIAS Part 2 Trial**

The ALIAS Part 2 Trial randomized its first subject in February 2009 and has enrolled approximately 225 subjects in the ensuing year. The ratio of thrombolysed to nonthrombolysed subjects currently exceeds 5:1. The DSMB has reviewed the safety data of the ALIAS Part 2 Trial 3 times since its initiation and has approved the continuation of the trial on each occasion. In addition, safety analyses based on
deaths within 30 days were conducted in December 2009 (based on first 15 deaths) and in April 2010 (based on the first 30 deaths), and there were no safety concerns based on the predefined guidelines specified in the Safety Monitoring Plan and Statistical Analysis Plan of the ALIAS Part 2 Trial. Thus, we are confident that the changes introduced in the ALIAS Part 2 Trial have resulted in an improved participant safety profile.

Discussion
In Part 1 of the ALIAS Trial, the 90-day death rate was greater in ALB- than in saline-treated subjects (Table 3; Figure), and this was chiefly accounted for by 15 excess deaths occurring on Days 5 to 30 postrandomization. This timing suggests that the excess deaths were not the direct consequence of ALB-associated volume expansion but rather were due to indirectly acting mechanisms. Although these deaths occurred predominantly in older subjects with large ischemic strokes who had received larger volumes of intravenous fluids in the first 48 hours (Table 4B), careful adjudication failed to identify specific additional factors contributing to deaths of individual ALB-treated subjects. Cardiovascular SAEs occurred more commonly after ALB administration than with saline, in particular, acute coronary syndrome and pulmonary edema, although the event rate of the latter was no greater than expected from the ALIAS Pilot Trial.13 In subjects with CV-SAEs, the clinical diagnosis of MI was more common in those receiving ALB (15 cases; 7 acute, 6 late) than in saline-treated subjects (2 acute or subacute cases); asymptomatic troponin elevations also tended to be more common in the former group. Nonetheless, deaths even in ALB-treated subjects with acute MI were much more commonly attributed to complications of a large stroke than to a direct cardiac mechanism (Table 3). These results, taken together, suggest that ALB treatment tended to predispose susceptible subjects to a degree of myocardial stress, which, acting indirectly and in combination with other predisposing factors, increased mortality in the 5- to 30-day period after treatment. This was supported by a comparison of overall mortality in the ALIAS Part 1 safety subjects who had not experienced a CV-SAE; with ALB, 33 deaths in 154 subjects (21.4%), and with saline, 35 deaths in 178 subjects (19.7%). That is, the entire difference in treatment-related death rates could be attributed to those ALB-treated subjects who experienced CV-SAEs.

The protocol modifications instituted in Part 2 were largely intended to diminish ALB-associated mortality in the very elderly and in subjects with fluid excess, and to reduce the likelihood of including subjects at higher risk of cardiovascular events. (To maximize the applicability of ALB treatment to ischemic stroke, we chose not to impose a ceiling on the permissible baseline NIHSS score.) Because this proposal involved major protocol modifications, the DSMB, NINDS, and Food and Drug Administration concurred in the decision that the study go forward from this point on as a separate trial—"the ALIAS Part 2 Trial"—and that the ALIAS Part 1 data eventually be used in a pooled analysis after completion of Part 2.

It is appropriate that we devote brief attention to the bioethical considerations that guided our decision-making as to publication. As laid forth in the Declaration of Helsinki,26 clinical investigators have an ethical duty to make publicly available the results of human subjects research, whether positive, negative, or inconclusive. The National Institutes of Health’s Belmont Report27 reaffirmed the concept of beneficence as a fundamental guiding principle of biomedical research.28 Randomized clinical trials tend to place doctors in the ethically challenging position of acting both as physicians and as scientists.29 In so doing, investigators must necessarily adopt a utilitarian approach that maximizes societal benefit at the same time as minimizing the risk to individual subjects. Thus, the challenge to the physician–investigator is to maintain clinical equipoise—a state of uncertainty about the relative merits of treatments A versus B—during his or her participation in a randomized controlled trial so as to avoid forming fixed beliefs about a novel treatment whose benefit has yet to be established.29,30

In publishing the safety results of the ALIAS Trial Part 1 at this time, we fulfill an ethical duty to publicize the results of our trial, but we are cognizant that these results may pose risks to clinical equipoise in the stroke–neurology community despite the fact that the DSMB, the clinical trials leadership at NINDS, and the Food and Drug Administration have all given their approval to continue the ALIAS Trial as Part 2. Baron31 has applied decision analysis to illustrate the implications of various decision-making scenarios. In our case, publication of the ALIAS Trial Part 1 data at this time poses the potential risk that certain centers might decide not to participate in Part 2 of the ALIAS Trial or to participate with reduced enthusiasm; and that publication might reinforce the beliefs of neuroprotection nihilists. By contrast, deferring publication might engender suspicion among participants despite our well-intentioned rationale. We have decided, on balance, that clinical equipoise is best maintained by publishing this report at this time.

The ALIAS Trial is a clinical trial that was adapted in midcourse. Because these changes were not preplanned, the trial does not meet the definition of "adaptive design."72 Nonetheless, we believe that describing the process that led to Part 2 may benefit other acute stroke clinical trials experiencing a similar predicament and thus needing to consider the potential incorporation of changes, particularly with respect to safety factors, into their study design to make it more efficient and truly adaptive.

Appendix
ALIAS Part 1 Executive Committee
Myron D. Ginsberg, University of Miami, Miami, Fla (Study Chair and Principal Investigator, Clinical Coordinating Center); Yoko Y. Palesch, Medical University of South Carolina, Charleston, SC (Principal Investigator, Statistics and Data Coordinating Center); Michael D. Hill, University of Calgary, Calgary, Canada (Director, Canadian Coordinating Center); Bonnie D. Waldman, Medical University of South Carolina, Charleston, SC (Project Manager); Lynn Patterson (ex officio), Medical University of South Carolina, Charleston, SC (Project Manager Assistant); Richard Leinster, Medical University of South Carolina, Charleston, SC (Data Manager); Isabel Mendez, University of Miami, Miami, Fla (Financial Man-
Enrolling Clinical Centers, ALIAS Part 1 Trial
(With Numbers of Subjects Contributed)
M. Hill, University of Calgary, Calgary, Canada (56); W. Clark, Oregon Health Sciences University, Portland, Ore (36); A. Shaub, University of Alberta, Edmonton, Canada (36); D. Selchen, Trillium Health Centre, Mississauga, Canada (35); M. Concha, Intercosta Neurology, Sarasota, Fla (15); J.-M. Boulanger, Hospital Charles LeMoyne, Greenfield Park, Canada (14); S. Silliman, University of Florida/Standlife, Jacksonville, Fla (13); A. Forteza, University of Miami, Miami, Fla (12); L. Pettigrew, University of Kentucky, Lexington, Ky (12); K. Remmel, University of Louisville, Louisville, Ky (10); M. Rymer, St Luke’s Hospital, Kansas City, Mo (10); V. Hachinski, London Health Sciences Centre, London, Canada (9); S. Hanson, Park Nicollet Institute, Minneapolis, Minn (9); N. Bayer, St Michael’s Hospital, Toronto, Canada (8); D. Chiu, Methodist Hospital, Houston, Texas (8); P. Teal, Vancouver Coastal Health Authority, Vancouver, Canada (8); R. Englehard, Sacred Heart Hospital, Eugene, Ore (7); L. Wechsler, University of Pittsburgh Medical Center, Pittsburgh, Pa (7); R. Dafer, Loyola University, Maywood, Ill (6); J. Gebel, Jewish Hospital, Louisville, Ky (6); G. Gubitz, Dalhousie University, Halifax, Canada (6); D. Laskowitz, Duke University, Durham, NC (6); S. Cruz-Flores, St Louis University, St Louis, Mo (5); G. Howell, Villages Research Group, Ocala, Fla (5); M. Jacoby, Ruan Neurology Clinic, Des Moines, Iowa (5); J. Schindler, Yale University, New Haven, Conn (5); R. Zweifler, University of South Alabama, Mobile, Ala (5); M. Beaudry, CSSS de Chicoutimi, Saguenay, Canada (4); C. Ionita, Millard Fillmore Gates Hospital, Buffalo, NY (4); G. Lopez, Baylor College of Medicine, Houston, Texas (4); S. Messe, University of Pennsylvania, Philadelphia, Pa (4); S. Sen, University of North Carolina, Chapel Hill, NC (4); R. Stephens, John Muir Medical Center, Walnut Creek and Concord, Calif (4); J. Teitelbaum, Montreal Neurological Institute, Montreal, Canada (4); C. Voll, University of Saskatchewan, Saskatoon, Canada (4); D. Camp, Seton Family of Hospitals, Austin, Texas (3); T. Collier, Royal Island Hospital, Kamloops, Canada (3); B. Coull, University of Arizona, Tucson, Ariz (3); M. Flaster, St Joseph’s Hospital, Phoenix, Ariz (3); R. Kelley, LSU Health Sciences Center, Shreveport, La (3); S. Mallenbaum, Neurological Consultants of Virginia Beach, Virginia Beach, Va (3); N. Solenski, University of Virginia, Charlottesville, Va (3); S. Starkman, UCLA, Los Angeles, Calif (3); G. Stotts, Ottawa Hospital, Ottawa, Canada (3); S. Bansil, Overlook Hospital, Summit, NJ (2); R. Fessler, St Joseph Mercy Oakland, Southfield, Mich (2); J. Hanna, MetroHealth Medical Center, Cleveland, Ohio (2); J. Harris, Neurological Consultants, Ft Lauderdale, Fla (2); H. Kirshner, Vanderbilt University, Nashville, Tenn (2); E. Leira, University of Iowa, Iowa City, Iowa (2); C. Lewandowski, Henry Ford Health System, Detroit, Mich (2); R. Welch, Wayne State University, Detroit, Mich (2); M. Aguilar, Mayo Clinic Hospital, Phoenix, Ariz (1); D. Brenner, University of Alabama, Birmingham, Ala (1); E. Feen, University Hospitals, Cleveland, Ohio (1); K. O’Phelan, Queens Medical Center, Honolulu, Hawaii (1); and D. Weissman, Abington Memorial Hospital, Abington, Pa (1).

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None.

References


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SUPPLEMENTAL MATERIAL

Consort diagram

Assessed for eligibility (n = 8761)

Excluded (n = 8327)
Not meeting inclusion criteria (n = 6206)
Refused to participate (n = 155)
Other reasons (n = 1966)

Randomized (n = 434)

Allocated to Albumin (n = 215)
  Received allocated intervention (n = 207)
  Did not receive allocated intervention (n = 8)

Allocated to Saline (n = 219)
  Received allocated intervention (n = 217)
  Did not receive allocated intervention (n = 2)

Completed the Study (n=141)
Not-Completed Reasons (safety cohort):
  Death (n=52)
  Lost to follow-up (n = 9)
  Consent withdrawn (n = 4)
  Administrative reason (n=1)

Completed the Study (n=174)
Not-Completed Reasons (safety cohort):
  Death (n=39)
  Lost to follow-up (n = 2)
  Consent withdrawn (n = 2)
  Administrative reason (n=0)

Analysis

Analyzed for Efficacy (n = 215)
Analyzed for Safety (n=207)

Analyzed for Efficacy (n = 219)
Analyzed for Safety (217)