Safety, Pharmacokinetics and Pharmacodynamics of DS-1040, in Combination with Thrombectomy, in Japanese Patients with Acute Ischemic Stroke

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Abstract

Background and Objectives DS-1040 is a novel inhibitor of the activated form of thrombin-activatable fibrinolysis inhibitor that may have therapeutic potential in thromboembolic diseases, such as acute ischemic stroke (AIS) or pulmonary embolism. We undertook a Phase I clinical trial to investigate the safety, pharmacokinetics, and pharmacodynamics of DS-1040 in Japanese patients who were eligible for thrombectomy following AIS.

Methods The trial enrolled patients with AIS due to large vessel occlusion, who were planned for thrombectomy within 8 h of symptom onset. Subjects were randomized to receive a single intravenous infusion of placebo or DS-1040 (0.6, 1.2, 2.4 or 4.8 mg) in a sequential-cohort design. The primary endpoints were the incidence of intracranial hemorrhage (ICH) and major extracranial bleeding within 36 and 96 h, respectively, of treatment initiation. Treatment-emergent adverse events (TEAEs) and pharmacokinetic/pharmacodynamic parameters were also assessed.

Results Nine patients received placebo and 32 patients received DS-1040. There were no cases of symptomatic ICH or major extracranial bleeding with either placebo or DS-1040 after 36 and 96 h. One patient, who received DS-1040 0.6 mg, experienced a subarachnoid hemorrhage that was considered to be drug-related. Three patients died (2 placebo, 1 DS-1040), but no deaths were adjudicated as study drug-related. In vivo exposure to DS-1040 increased in proportion to dosage, but no clear dose-response relationship was seen for D-dimer levels and thrombin-activatable fibrinolysis inhibitor activity.

Conclusions Single doses of DS-1040 0.6–4.8 mg were well tolerated in Japanese patients with AIS undergoing thrombectomy.

Clinical trial registration number NCT03198715; JapicCTI-163164.

1 Introduction

Acute ischemic stroke (AIS) is a medical emergency and a major cause of death and disability worldwide. In Japan, approximately 195,000 cases of AIS occur each year, and there were over 90,000 deaths due to the disease in 2019 [1].

Stroke-related mortality is declining because of general improvements in the acute management of AIS, including the development of specialized stroke centers and the availability of effective revascularization with thrombolytic drugs and endovascular therapy (thrombectomy) [2]. In a meta-analysis of five randomized controlled trials, thrombectomy was associated with a reduced risk of disability: 46% of patients who received thrombectomy had a favorable outcome (i.e., a modified Rankin Scale [mRS] score of 0–2 at 90 days), versus 26.5% of control patients ($p < 0.0001$) [3].

Higher rates of recanalization and the prevention of reocclusion are considered to be significant factors in improving outcomes in AIS [4–6], and the number of patients treated with thrombectomy is increasing. However, more needs to be done to improve outcomes in AIS, with data showing variable rates of recanalization and effects on prognosis [7]. For example, in the SWIFT PRIME trial, thrombectomy (using a stent retriever device) plus alteplase...
Key Points

DS-1040, a small-molecule inhibitor of the activated form of thrombin-activatable fibrinolysis inhibitor, has fibrinolytic activity and may improve reperfusion rates and prognostic indicators in patients with acute ischemic stroke (AIS).

We conducted a Phase I study to assess the safety/tolerability and pharmacokinetics/pharmacodynamics of DS-1040 in Japanese patients with AIS who were eligible for thrombectomy.

Single intravenous doses of DS-1040 (0.6–4.8 mg) were safe and well tolerated. There was limited evidence of a dose-effect relationship or marked differences between DS-1040 and placebo on pharmacodynamic parameters or scales measuring stroke severity or associated disability and, thus, further research is required to fully investigate the effects of DS-1040 in patients with AIS.

DS-1040-A-J110 was a randomized, placebo-controlled, single-blind, dose-escalation trial undertaken to investigate the safety, pharmacokinetics, and pharmacodynamics of DS-1040 in patients with AIS who were eligible for thrombectomy (NCT03198715; JapicCTI-173612). The trial was performed at 20 centers in Japan (Supplementary Table S1; Online Resource 1) between July 2017 and January 2020 and was conducted in compliance with the Declaration of Helsinki, the International Conference on Harmonization Integrated Guideline for Good Clinical Practice, and additional regulatory codes applicable in Japan. The trial protocol was approved by the Institutional Review Board at each participating center before any patients were enrolled at that center. All patients, or a legally competent representative, gave written informed consent to participate.

2.1 Inclusion and Exclusion Criteria

Complete inclusion and exclusion criteria are listed in Supplementary Table S2 (Online Resource 1).

AIS patients aged 20–89 years were screened for enrollment. To be included, patients had to have angiogram-confirmed (by computed tomography or magnetic resonance imaging) internal carotid artery (ICA) or middle cerebral artery (MCA; M1 region) occlusion, for which thrombectomy was planned as the primary treatment modality. Only those considered able to receive thrombectomy within 8 h of the onset of ischemic symptoms were included. Additionally, patients had to have a National Institutes of Health Stroke Scale (NIHSS) score of ≥ 6 and an Alberta Stroke Program Early CT Score of ≥ 6.

Patients were excluded if they: (i) were planned for, or had received, thrombolytic therapy; (ii) had evidence of (or risk factors for) hemorrhagic stroke, or other major bleeding event; (iii) had severe renal or hepatic impairment; or (iv) had a contraindication to thrombectomy. DS-1040 was not used in combination with thrombolytic therapy because of the potential for bleeding events.

2.2 Study Design

The study design is shown in Fig. 1. Patients were enrolled in four sequential cohorts. Cohort 1 consisted of six patients, all of whom received a single dose of DS-1040 0.6 mg. The study design allowed for a further six patients to be recruited into this cohort if study discontinuation or interruption criteria were met (Supplementary Table S3; Online Resource 1). On the basis of previous modeling and simulation work in healthy volunteers, we calculated that a DS-1040 dosage of 1.2 mg would achieve maximum plasma drug concentrations ($C_{\text{max}}$) of 23.8–31.7 ng/mL, versus a reported 90% of the maximal inhibitory concentration (IC$_{90}$) of 38.2 ng/mL in clot lysis assays [10]. We adopted a conservative approach and selected 0.6 mg as the starting dose, due to the lack of safety data in the Japanese population. Cohorts 2, 3 and 4 each consisted of 16 patients, randomized by...
block assignment to the DS-1040 (1.2 mg, 2.4 mg, or 4.8 mg, respectively) or placebo groups in a 3:1 ratio. Thus, the total planned enrollment was 54 or 60 patients, depending on the outcome of Cohort 1.

Enrollment into Cohorts 2, 3 and 4 began once all patients in the previous cohort had completed 96 h of follow-up, and an evaluation of the available safety data at that time-point supported the continuation of the trial. Within each cohort, patients were evaluated at 36 h for the presence of symptomatic intracerebral hemorrhage (ICH) before the next patient was enrolled. This continued until three patients aged ≥ 81 years (considered to be at higher risk of bleeding events) had been randomized to receive DS-1040 in each cohort.

2.3 Treatment

Both DS-1040 and placebo were administered by intravenous (IV) infusion over 6 h. Each dose of DS-1040 was added to 100 mL normal saline for infusion; placebo infusions consisted of 100 mL normal saline only. The first 25% of each dose was administered over 30 min, and the remaining 75% over 5.5 h. DS-1040 was supplied to each study center by Daiichi Sankyo Co., Ltd. (Tokyo, Japan). Placebo supplies were taken from IV fluid stocks at each center.

Study treatment was started within 8 h of symptom onset, prior to thrombectomy or as soon as possible afterwards. Except for monotherapy with aspirin (acetylsalicylic acid), clopidogrel or enoxaparin, and heparin used in connection with thrombectomy, drugs with thrombolytic, antiplatelet, or anticoagulant activity were prohibited until 24 h after the start of study treatment.

2.4 Outcomes

Patients were followed up for 90 days from the date of study treatment. The main objective of the trial was to assess the safety of DS-1040; accordingly, the primary endpoints were: (i) the incidence of symptomatic (according to the European Cooperative Acute Stroke Study [ECASS] criteria) and asymptomatic ICH within 36 h of starting study treatment; (ii) TIMI major extracranial bleeding; (iii) Bleeding events / adverse events / serious adverse events.
and (ii) extracranial (Thrombolysis in Myocardial Infarction [TIMI]) major bleeding within 96 h of starting study treatment. The ECASS definition of symptomatic ICH is ICH with clinical deterioration causing an increase in the NIHSS score of ≥ 4 points, and the definition of TIMI major bleeding is clinically evident bleeding with a 5 g/dL or greater decrease in hemoglobin; a 2-unit blood transfusion (where 1 unit is equivalent to 200 mL) was converted to a 1 g/dL increase in hemoglobin. The final classification and assessment of hemorrhagic events was performed by a central adjudication committee member who was blinded to the patient's treatment allocation.

Secondary endpoints were: (i) treatment-emergent adverse events (TEAEs); (ii) pharmacokinetic parameters, calculated using a non-compartmental model and DS-1040 concentrations measured in plasma; (iii) pharmacodynamic parameters (D-dimer levels and TAFIa activity); (iv) change in NIHSS score from baseline to 24 (± 4) h after the start of study treatment; and (v) the proportion of patients with a favorable outcome (defined as a mRS score of 0, 1, or 2) at 30 and 90 days. Pharmacokinetic parameters and dose proportionality were calculated using WinNonlin™ version 6.0 or higher (Certara, Princeton, NJ, USA).

As an exploratory endpoint, we investigated the proportion of patients achieving complete (i.e., modified Thrombolysis in Cerebral Infarction [TICI] grade 3 [12]) reperfusion after thrombectomy, and the impact of DS-1040 on the proportion of patients with TICI grade 3.

### 2.4.1 Adverse Events

Information on AEs was collected at all study visits, including the day of study drug administration (Visit 1). AEs were defined as untoward medical occurrences, without regard to causality. An AE was considered to be serious if: (i) it was life-threatening; (ii) it was fatal; (iii) it resulted in hospitalization or prolongation of existing hospitalization; (iv) it resulted in persistent or significant disability or incapacity; or (iii) it was otherwise considered important. TEAEs were defined as AEs that were not present prior to the study but developed between the start of study treatment and Day 30 (Visit 5).

### 2.4.2 Bioanalytical Assay

Blood samples for pharmacokinetic analysis were taken before the start of study drug administration, and at 0.5, 3, 6, 18, 24, 48 and 96 h after the start of treatment. For measurement of plasma DS-1040 concentrations, 3 mL of venous or arterial blood from each subject was collected in K2-EDTA tubes. After blood collection, tubes were mixed by inversion immediately and then cooled with ice. Plasma was obtained by centrifugation (4 °C, 1500 × g, 15 min) and was stored at or below − 20 °C until analysis.

The assay method was validated for a range of concentrations between 0.500 and 500 ng/mL, based on the analysis of 0.100 mL of plasma. DS-1040 and its internal standard were extracted from plasma samples (0.100 mL) through solid phase extraction using Strata® SCX (55 µm, 70 Å), 50 mg/well solid phase extraction plates (Phenomenex, Torrance, CA, USA). The final eluent was evaporated to dryness, and the reconstituted extract was injected into a Sciex API 4000™ (AB Sciex, Framingham, MA, USA) tandem mass spectrometer with turbo ion spray source by electrospray ionization in the positive ion mode. Multiple-reaction monitoring of DS-1040 (m/z 294→180) and its internal standard DS-1040-D4 (m/z 298→184) was conducted using an HPLC column. Chromatographic separation was performed using a Luna C18(2) 3 µm, 100 × 2.0 mm column (Phenomenex). The assay accuracy was in the range − 6.4% to − 1.3%, and the lower limit of quantification was 0.5 ng/mL.

Blood samples for pharmacodynamic analysis were taken before the start of study drug administration and at 6, 24 and 48 h afterwards. On each occasion, 9 mL of venous blood was collected in sodium citrate 3.2% tubes, which were immediately mixed by inversion. D-dimer was measured using the STA®-Liatest® D-Di assay, while TAFIa activity was measured using the STA®-Stachrom® TAFI assay (both manufactured by Diagnostica Stago S.A.S., Asnières sur Seine, France).

D-dimer levels were reported in µg/mL fibrinogen equivalent units (FEU), over an analytical measuring range (AMR) of 0.36–4.00 µg/mL FEU. Samples with results below the AMR were reported as < 0.36 µg/mL FEU; those with results > 4.00 µg/mL FEU were re-assayed after diluting up to 5-fold using Owren-Koller buffer, and the results multiplied by the corresponding dilution factor. The reference range for D-dimer in citrated plasma is < 0.50 µg/mL FEU. TAFIa was measured and reported as a percentage, with an AMR of 9–137% and a reference range of 77–177%. Samples with results below the AMR were reported as < 9%, and those > 137% were re-assayed after diluting up to 8-fold using Owren-Koller buffer.

### 2.5 Statistical Analysis

Safety analyses were performed on the safety analysis set, defined as the population of enrolled patients who received at least one dose of study treatment. For all endpoints, including the primary endpoints, point estimates were calculated for each dosage of DS-1040, and for all placebo recipients as a single group. For TEAEs, incidences were calculated for each treatment group. Pharmacokinetic and pharmacodynamic parameters were assessed for all patients without major protocol violations and with at least one
plasma concentration or pharmacodynamic measurement, respectively. Summary statistics on plasma concentrations, pharmacokinetic parameters and pharmacodynamic measurements were calculated for each treatment group.

Measures of efficacy (i.e., change in NIHSS score from baseline to 24 h; proportion with favorable outcome at 30 and 90 days) were assessed for the full analysis set, which included all patients who received study treatment and for whom relevant NIHSS or mRS data were available. Summary statistics were calculated for efficacy endpoints.

All statistical analyses were performed using SAS® version 9.4 or higher (SAS Institute, Cary, NC, USA).

3 Results

3.1 Patient Disposition

Enrollment into DS1040-A-J110 was stopped while the trial was in progress because the sponsor decided to discontinue the late-stage development of DS-1040.1 At this point, Cohorts 1, 2, and 3 had been fully recruited. Four of the planned 16 patients had been enrolled in Cohort 4: three patients were randomized to DS-1040 4.8 mg and one patient was randomized to placebo.

Forty-two patients were randomized (33 to DS-1040 and 9 to placebo), and only one patient (in the DS-1040 2.4 mg group) did not receive study drug (Supplementary Figure S1; Online Resource 1). Thus, the safety analysis set and full analysis set both included 41 patients.

No events occurred that would have triggered the recruitment of additional patients into Cohort 1. Recruitment into Cohort 2 was therefore able to begin once the first six patients in Cohort 1 had completed 96 h of follow-up.

Five patients discontinued the study. Discontinuation was due to death in three patients (one patient who received DS-1040 2.4 mg, and two patients who received placebo), none of which were judged to be related to study drug. Additionally, one patient in the 1.2 mg group was withdrawn from the study because of a suspected protocol violation relating to the exclusion criteria; another patient (in the 2.4-mg group; already described above) was withdrawn before treatment started because angiography suggested a risk of hemorrhage.

3.2 Patient Characteristics

There were no important differences between the study groups at baseline (Table 1). Patients in the safety analysis set had a mean (± standard deviation [SD]) age of 74.5 ± 9.0 (range 53.0–87.0) years, and approximately half (46.3%) were female. NIHSS scores ranged from 6 to 33, and median scores were similar for DS-1040 and placebo recipients (20.0 vs 21.5, respectively).

3.3 Primary Endpoints

Rates of ICH and major extracranial bleeding are shown in Table 2. There were no cases of symptomatic ICH in the 36 h after the start of treatment among either DS-1040 or placebo recipients. Similarly, no cases of major extracranial bleeding were reported in the first 96 h. The incidence of asymptomatic ICH was 37.5% among DS-1040 recipients and 11.1% among placebo recipients; no relationship between DS-1040 dose and incidence of asymptomatic ICH was seen.

3.4 Secondary Endpoints

3.4.1 Adverse Events

TEAEs are summarized in Table 3. Overall, DS-1040 was well tolerated: although most patients (35/41 patients; 85.4%) experienced AEs, only one TEAE (a subarachnoid hemorrhage in a patient who received DS-1040 0.6 mg) was considered to be study drug–related and was not categorized as serious. Overall, TEAEs classed as serious occurred infrequently (reported in five [12.2%] patients), and none were considered related to study drug. Furthermore, there were no serious bleeding events, and no discontinuations of study drug due to a TEAE in any treatment group.

Among DS-1040 recipients, there was no clear evidence of a dose-proportional increase in TEAEs (Table 3). The most frequent TEAEs (i.e., those reported in > 10% [i.e., > 3] of DS-1040 recipients) were constipation (25.0% vs 33.3% with placebo), hemorrhagic infarction (25.0% vs 0.0%), skin exfoliation (15.6% vs 0.0%) and pneumonia aspiration (12.5% vs 22.2%).

Three patients died during the trial; all deaths were considered unrelated to study drug. One patient who received DS-1040 2.4 mg had an acute myocardial infarction 9 days after study drug administration. The other two deaths were in placebo recipients; these were due to aspiration pneumonia in one patient and an exacerbation of right lung adenocarcinoma in the other.

1 The addition of DS-1040 to enoxaparin did not result in improved efficacy compared with enoxaparin alone in a Phase II trial in patients with acute pulmonary embolism (NCT02923115) [13].
3.4.2 Pharmacokinetics

The plasma concentration-time curves for each dosage of DS-1040 are shown in Fig. 2, and calculated pharmacokinetic parameters are summarized in Table 4. In vivo exposure to DS-1040 (measured as the area under the plasma concentration-time curve [AUC]) increased in a dose-proportional manner, as did \( C_{\text{max}} \). As expected, mean time to \( C_{\text{max}} \) increased with dosage, from 2.21 h at a dosage of 0.6 mg to 5.96 h at a dosage of 4.8 mg. The mean elimination half-life (\( t_{\frac{1}{2}} \)) and volume of distribution at steady state (\( V_{\text{ss}} \)) of DS-1040 also showed dose-dependency, increasing from

\[ \Delta \text{Adis} \]
Table 3  Treatment-emergent adverse events (TEAEs). In the analysis of TEAEs by preferred term, only those TEAEs that occurred in at least 2 patients in any treatment group (i.e. with placebo, with any individual dosage of DS-1040, or with DS-1040 overall) are included, and are listed in descending order by system organ class for DS-1040 recipients (i.e. the second-to-last column from the right)

| Type of adverse event (AE), n (%) | Placebo (n = 9) | DS-1040 (n = 32) | Overall (n = 41) |
|----------------------------------|----------------|------------------|-----------------|
|                                  |                | 0.6 mg (n = 6)   | 1.2 mg (n = 12) | 2.4 mg (n = 11) | 4.8 mg (n = 3) | Total (n = 32) |
| **Overall results**               |                |                  |                 |                 |                 |                |
| Any AE                           | 8 (88.9)       | 5 (83.3)         | 10 (83.3)       | 10 (90.9)       | 2 (66.7)        | 27 (84.4)      | 35 (85.4)      |
| TEAE                             | 8 (88.9)       | 5 (83.3)         | 10 (83.3)       | 10 (90.9)       | 2 (66.7)        | 27 (84.4)      | 35 (85.4)      |
| Drug-related TEAE                | 0 (0.0)        | 1 (16.7)         | 0 (0.0)         | 0 (0.0)         | 0 (0.0)         | 1 (3.1)        | 1 (2.4)        |
| Bleeding TEAE<sup>a</sup>         | 6 (66.7)       | 3 (50.0)         | 6 (50.0)        | 9 (81.8)        | 2 (66.7)        | 20 (62.5)      | 26 (63.4)      |
| Drug-related bleeding TEAE<sup>a</sup> | 0 (0.0)   | 1 (16.7)         | 0 (0.0)         | 0 (0.0)         | 0 (0.0)         | 1 (3.1)        | 1 (2.4)        |
| Serious TEAE                     | 1 (11.1)       | 0 (0.0)          | 2 (16.7)        | 2 (18.2)        | 0 (0.0)         | 4 (12.5)       | 5 (12.2)       |
| **Analysis by system organ class and preferred term** | | | | | | |
| Gastrointestinal disorders       |                |                  |                 |                 |                 |                |
| Constipation                     | 3 (33.3)       | 2 (33.3)         | 2 (16.7)        | 3 (27.3)        | 1 (33.3)        | 8 (25.0)       | 11 (26.8)      |
| Gingival bleeding                | 1 (11.1)       | 0 (0.0)          | 1 (8.3)         | 1 (9.1)         | 0 (0.0)         | 2 (6.3)        | 3 (7.3)        |
| Vomiting                         | 1 (11.1)       | 2 (33.3)         | 0 (0.0)         | 0 (0.0)         | 0 (0.0)         | 2 (6.3)        | 3 (7.3)        |
| Nervous system disorders         |                |                  |                 |                 |                 |                |
| Subarachnoid hemorrhage          | 1 (11.1)       | 1 (16.7)         | 1 (8.3)         | 1 (9.1)         | 0 (0.0)         | 3 (9.4)        | 4 (9.8)        |
| Hemorrhagic cerebral infarction  | 0 (0.0)        | 0 (0.0)          | 1 (8.3)         | 1 (9.1)         | 1 (33.3)        | 3 (9.4)        | 3 (7.3)        |
| Skin and subcutaneous tissue disorders | | | | | | |
| Dermatitis diaper                | 2 (22.2)       | 1 (16.7)         | 2 (16.7)        | 0 (0.0)         | 0 (0.0)         | 3 (9.4)        | 5 (12.2)       |
| Skin exfoliation                 | 0 (0.0)        | 1 (16.7)         | 3 (25.0)        | 1 (9.1)         | 0 (0.0)         | 5 (15.6)       | 5 (12.2)       |
| Vascular disorders               |                |                  |                 |                 |                 |                |
| Hemorrhagic infarction           | 0 (0.0)        | 1 (16.7)         | 5 (41.7)        | 2 (18.2)        | 0 (0.0)         | 8 (25.0)       | 8 (19.5)       |
| Hemorrhage                       | 0 (0.0)        | 0 (0.0)          | 0 (0.0)         | 2 (18.2)        | 0 (0.0)         | 2 (6.3)        | 2 (4.9)        |
| Renal and urinary disorders      |                |                  |                 |                 |                 |                |
| Hemorrhage urinary tract         | 1 (11.1)       | 0 (0.0)          | 1 (8.3)         | 2 (18.2)        | 0 (0.0)         | 3 (9.4)        | 4 (9.8)        |
| Hematuria                        | 1 (11.1)       | 1 (16.7)         | 0 (0.0)         | 1 (9.1)         | 0 (0.0)         | 2 (6.3)        | 3 (7.3)        |
| Respiratory, thoracic, and mediastinal disorders | | | | | | |
| Pneumonia aspiration             | 2 (22.2)       | 1 (16.7)         | 2 (16.7)        | 1 (9.1)         | 0 (0.0)         | 4 (12.5)       | 6 (14.6)       |
| General disorders and administration site conditions | | | | | | |
| Pyrexia                          | 1 (11.1)       | 1 (16.7)         | 0 (0.0)         | 2 (18.2)        | 0 (0.0)         | 3 (9.4)        | 4 (9.8)        |
| Vessel puncture site hematoma    | 0 (0.0)        | 1 (16.7)         | 1 (8.3)         | 0 (0.0)         | 0 (0.0)         | 2 (6.3)        | 2 (4.9)        |
| Infections and infestations      |                |                  |                 |                 |                 |                |
| Urinary tract infection          | 2 (22.2)       | 0 (0.0)          | 1 (8.3)         | 1 (9.1)         | 0 (0.0)         | 2 (6.3)        | 4 (9.8)        |
| Respiratory tract infection      | 2 (22.2)       | 0 (0.0)          | 0 (0.0)         | 0 (0.0)         | 0 (0.0)         | 0 (0.0)        | 2 (4.9)        |
| Injury, poisoning, and procedural complications | | | | | | |
| Subcutaneous hematoma            | 2 (22.2)       | 0 (0.0)          | 1 (8.3)         | 1 (9.1)         | 1 (33.3)        | 3 (9.4)        | 5 (12.2)       |
| Psychiatric disorders            |                |                  |                 |                 |                 |                |
| Insomnia                         | 0 (0.0)        | 0 (0.0)          | 0 (0.0)         | 2 (18.2)        | 1 (33.3)        | 3 (9.4)        | 3 (7.3)        |
| Delirium                         | 0 (0.0)        | 0 (0.0)          | 2 (16.7)        | 0 (0.0)         | 0 (0.0)         | 2 (6.3)        | 2 (4.9)        |
| Metabolism and nutrition disorders |            |                  |                 |                 |                 |                |
| Hypokalemia                      | 0 (0.0)        | 0 (0.0)          | 2 (16.7)        | 0 (0.0)         | 0 (0.0)         | 2 (6.3)        | 2 (4.9)        |

<sup>a</sup>As judged by a member of the central adjudication committee
4.61 h and 17.6 L, respectively, in Cohort 1 (n = 1) to 29.7 h and 46.5 L, respectively, in Cohort 4 (n = 3).

3.4.3 Pharmacodynamics

D-dimer levels and TAFIa activity are shown in Fig. 3. At baseline, mean ± SD D-dimer levels were higher and showed greater variance in patients who received placebo than in those who received DS-1040 (3.1 ± 6.8 vs 1.3 ± 2.8 μg/mL FEU), but TAFIa activity was similar in both groups.

D-dimer levels increased by similar amounts between baseline and 24 h in the placebo and DS-1040 groups (mean change +0.28 vs +0.35 μg/mL FEU, respectively). Furthermore, mean increases in D-dimer levels were smaller in patients who received DS-1040 4.8 mg than in those who received 2.4 mg (+0.45 vs +1.18 μg/mL FEU at 24 h and +0.67 vs +1.03 μg/mL FEU at 48 h). However, consistent with its mechanism of action, doses of DS-1040 above 1.2 mg were associated with decreases in TAFIa activity at 6 h. There was evidence of a dose-response relationship at 6 h, since greater decreases in activity were seen with DS-1040 4.8 mg than at lower doses, but this difference tended to disappear at later time-points.

3.4.4 NIHSS and mRS Scores

Mean changes in NIHSS score between baseline and 24 h are shown in Fig. 4. Although DS-1040 was associated

with reductions in NIHSS score at all dose levels, there was no strong evidence of dose proportionality; furthermore, the magnitude of change was comparable for placebo and DS-1040 doses up to 2.4 mg.

Among DS-1040 recipients, the proportion of patients with a favorable outcome (mRS score 0–2) was 43.8% (95% confidence interval [CI]: 26.4–62.3%) and 53.1% (95% CI 34.7–70.9%) at 30 and 90 days, respectively. Values for placebo were 11.1% (95% CI 0.3–48.2%) at 30 days and 33.3% (95% CI 7.5–70.1%) at 90 days. At both time-points, the proportion of patients with a favorable outcome was higher for DS-1040 than for placebo, regardless of DS-1040 dosage (Fig. 5). However, there was no apparent relationship between DS-1040 dosage and the probability of favorable outcome at either 30 or 90 days.

3.5 Exploratory Endpoints

3.5.1 TICI Grade

TICI grade score was available for 40 patients (31 in the DS-1040 group and 9 in the placebo group). Except for one patient who received DS-1040 1.2 mg, all patients achieved either TICI grade 2B or grade 3 reperfusion after thrombectomy. The percentage of patients with complete (TICI grade
3) reperfusion was 44.4% (n = 4) in the placebo group and 51.6% (n = 16) in the DS-1040 group. There was no apparent relationship between DS-1040 treatment (or dosage) and the probability of achieving TICI grade 3 reperfusion.

4 Discussion

DS1040-A-J110 was the first clinical trial of DS-1040 in patients with AIS in Japan, and, indeed, one of the first studies of DS-1040 in any indication. Accordingly, its primary objective was to evaluate the safety of DS-1040, with the primary endpoints being the incidence of symptomatic or asymptomatic ICH or major extracranial bleeding. Over a range of single doses of DS-1040 from 0.6 to 4.8 mg, given by IV infusion over 6 h, we found no evidence of symptomatic ICH in the first 36 h after the start of treatment, and no major extracranial bleeding in the first 96 h. In the current study, asymptomatic ICH was more common in patients receiving DS-1040 than placebo (37.5% vs 11.1%) but had no impact on the outcome of treatment. In a sub-analysis of data from the RESCUE-Japan registry, the incidence of asymptomatic ICH among Japanese patients treated with thrombectomy for large vessel occlusion was 26.4%, compared with 13.7% in patients who received conservative therapy or thrombolysis (p < 0.001) [14]. In the current study, the incidence of asymptomatic ICH in the placebo group tended to be lower than in the RESCUE-Japan registry.

Thus, our study confirms the conclusions of earlier work [9–11], which suggested that the potential of DS-1040 to cause clinically significant bleeding events is low. Additionally, DS-1040 was well tolerated, with drug-related TEAEs occurring in only one DS-1040–treated patient. Bleeding events, including major bleeding, occurred at a similar frequency in the placebo and DS-1040 groups, and there were no serious drug-related TEAEs.

Table 4  Pharmacokinetic parameters of DS-1040

| Cohort | Dosage (mg) | n\(^a\) | \(C_{\text{max}}\) (ng/mL) | \(t_{\text{max}}\) (h) | \(AUC_{0-24\text{h}}\) (ng × h/mL) | \(AUC_{\text{last}}\) (ng × h/mL) | \(t_{1/2}\) (h) | CL (L/h) | \(V_{\text{ss}}\) (L) |
|--------|------------|--------|-----------------|--------|-----------------|-----------------|--------|---------|-----------|
| 1      | 0.6        | 6      | 17.2 ± 6.25    | 2.21 ± 2.18 | 151 ± 40.5    | 111 ± 45.5     | 4.61\(^b\) | 3.39\(^b\) | 17.6\(^b\) |
| 2      | 1.2        | 11     | 29.7 ± 7.50    | 4.67 ± 2.29 | 267 ± 61.6    | 275 ± 72.2     | 16.8 ± 1.74\(^c\) | 3.03 ± 0.23\(^c\) | 22.6 ± 2.43\(^c\) |
| 3      | 2.4        | 10     | 60.6 ± 11.8    | 5.11 ± 1.85 | 503 ± 154     | 570 ± 230      | 19.5 ± 10.5\(^d\) | 4.62 ± 1.85\(^d\) | 43.3 ± 24.4\(^d\) |
| 4      | 4.8        | 3      | 141.0 ± 61.7   | 5.96 ± 0.03 | 1320 ± 752    | 1550 ± 962     | 29.7 ± 2.35 | 3.71 ± 1.73 | 46.5 ± 16.3 |

All data are expressed as mean ± standard deviation except where n = 1

\(AUC\) area under the plasma concentration-time curve, \(AUC_{0-24\text{h}}\) AUC from zero to 24 h, \(AUC_{\text{last}}\) AUC from zero to last measurable concentration, CL clearance, \(C_{\text{max}}\) maximum plasma concentration, n number of subjects, \(t_{1/2}\) elimination half-life, \(t_{\text{max}}\) time to \(C_{\text{max}}\), \(V_{\text{ss}}\) volume of distribution at steady state

\(^a\)Except where otherwise indicated
\(^b\)n = 1
\(^c\)n = 3
\(^d\)n = 5

Fig. 3  Effects of DS-1040 and placebo on A D-dimer levels, and B TAFI\(^a\) activity. Data points and error bars represent the mean and standard deviation, respectively. FEU fibrinogen equivalent units, NPP normal pooled plasma, TAFI thrombin-activatable fibrinogen inhibitor, TAFI\(^a\) activated form of TAFI
Alteplase and thrombectomy are standard treatments for AIS and should ideally be used in combination in eligible patients. Alteplase remains a standard of care for patients with radiologically confirmed AIS, provided that it can be administered within 9 h of symptom onset and that there are no contraindications to its use [15, 16]. However, the recanalization rate after alteplase is only approximately 40% in patients with distal MCA occlusion and < 10% in those with ICA occlusion [17]. In accordance with international and Japanese guidelines, thrombectomy is suitable only when a major artery is occluded [18–20].

There is undoubtedly a need for new additive or substitutive approaches, pharmacologic and otherwise, to the management of AIS with the potential to preserve brain tissue, further reduce mortality, and prevent disability resulting from neurologic damage. The discovery of TAFI and its activated form, TAFIa, which acts as a physiologic ‘brake’ on plasmin-mediated fibrinolysis [21], led to the development of several novel TAFIa inhibitors that demonstrated fibrinolytic activity in animal models and early clinical trials [22–25]. However, to date, none of these agents has been made commercially available.

The effects of DS-1040 on pharmacodynamic parameters and on NIHSS and mRS scores are challenging to interpret, with limited evidence of a dose-effect relationship or marked differences between DS-1040 and placebo. Although TAFIa activity levels decreased in DS-1040 recipients at one or more time-points, consistent, dose-related increases in D-dimer were not observed. Because blood clots in the brain tend to be small, changes in D-dimer levels may be too subtle to detect in peripheral blood.

It could be suggested that, in general, dose-proportionality was not observed because of the small number of patients in the highest dosage group, and because the dosages of DS-1040 used were lower than in previous studies [10, 11]. However, these previous studies, which used a wider range of doses, also failed to demonstrate dose-proportional effects of DS-1040 on D-dimer levels [10, 11]. In contrast, TAFIa activity levels have been shown to be reduced by DS-1040 in a dose-proportional manner [9].

**Fig. 4** Mean change in NIHSS score from baseline to 24 h. Error bars indicate standard deviation. NIHSS National Institutes of Health Stroke Scale

**Fig. 5** Modified Rankin Scale scores at A 30 and B 90 days after the start of treatment. mRS modified Rankin Scale
There was also no clear relationship between DS-1040 dosage and change in either the NIHSS score or the proportion of patients with a favorable outcome (mRS score 0–2). All but one patient (who received DS-1040 1.2 mg) in the overall study population achieved successful recanalization (TICI2b-3) with thrombectomy; however, a favorable outcome (mRS score 0–2) was seen in ~25% to 100% of patients in the DS-1040 arms compared with only 11% of patients in the placebo arm, despite successful recanalization. Although DS-1040 4.8 mg performed better than placebo with respect to these measures, no firm conclusions can be drawn because only three patients received DS-1040 4.8 mg.

Strict adherence to a trial protocol, particularly with respect to enrollment, drug administration and blood sampling, can be very challenging in the context of a medical emergency such as AIS. Therefore, a strength of our study was the clinical trial implementation system that we developed for use at each medical institution, since communication and cooperation between the sub-principal investigator, clinical research coordinator, and clinical trial staff was considered critical to ensure the validity and robustness of the data obtained.

We acknowledge the limitations of our trial, including the small number of patients included in each cohort; this precluded formal statistical analysis. Consequently, no firm conclusions can be drawn with regard to differences between DS-1040 doses, or between DS-1040 and placebo, for any of the secondary endpoints. Furthermore, our study was conducted exclusively in Japanese patients, and so the results may not be generalizable to other populations. A Phase Ib/II placebo-controlled trial (clinicaltrials.gov identifier: NCT02586233) of DS-1040 in patients with AIS has been conducted in Europe, North America, South Korea, Taiwan, and Australia. The preliminary results of this trial are available online and are expected to be published in due course.

5 Conclusions

In this Phase I study, single IV doses of DS-1040 0.6–4.8 mg were safe and well tolerated in Japanese patients with AIS undergoing thrombectomy. Further clinical research is needed to fully investigate the efficacy and safety of DS-1040 in a wider range of patients with AIS.

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Declarations

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Conflicts of Interest Nobuyuki Sakai received honoraria from Daiichi Sankyo in relation to the organization of this study and for non-related work, and non-related honoraria from Asahi-Intec, Biomedical Solutions, Medtronic, Stryker, and Terumo. Masataka Takeuchi reports the receipt of lecture fees from Stryker. Hirotsoshi Imamura has received non-related honoraria from Medtronic and Stryker. Shinichi Yoshimu- ra has received speakers’ bureau fees or honoraria from Boehringer Ingelheim, Daiichi Sankyo, Bayer, Bristol-Myers Squibb, Otsuka Pharmaceutical, Stryker, Kaneka Medics and Medtronic. Kazunori Toyoda reports personal fees from Daiichi Sankyo, Bayer Yakuhin, Bristol-Myers Squibb, Takeda and Nippon Boehringer-Ingelheim, unrelated to the submitted work. Hiroshi Yamagami reports receipt of research grants from Bristol-Myers Squibb; lecture fees from Stryker, Terumo, Medtronic, Johnson and Johnson, Bayer, Daiichi Sankyo, Bristol-Myers Squibb, Boehringer Ingelheim and Otsuka Pharmaceutical; and membership of advisory boards for Daiichi Sankyo in a non-related trial. Shigeru Fujimoto has received lecture fees from Takeda, Nippon Boehringer Ingelheim, Dai-Nippon Sumitomo Pharma, Bayer Yakuhin, Otsuka Pharmaceutical, Pfizer Japan, Daiichi Sankyo, Eisai, Bristol-Myers Squibb, Sanofi K.K. and MSD K. K., and received an honorarium from Daiichi Sankyo in relation to the organization of this study. Norihito Shimamura, Hiroichi Naito, Naoto Kimura, Osamu Masuo, Nobuyuki Hirotsune, Kenichi Morita, and Hideyuki Iishihara have no conflicts of interest to declare. Takafumi Nakatsu, Naoki Miyoshi and Miharu Suda are employees of Daiichi Sankyo.

Ethics approval The trial was performed in compliance with the Declaration of Helsinki, the International Conference on Harmonization Integrated Guideline for Good Clinical Practice, and additional regulatory codes applicable in Japan. The trial protocol was approved by the Institutional Review Board at each participating center before any patients were enrolled at that center.

Consent to participate All patients, or a legally competent representative, gave written informed consent to participate.

Consent for publication Not applicable.

Availability of data and material De-identified individual participant data (IPD) and applicable supporting clinical trial documents (study protocol, statistical analysis plan, informed consent form, clinical study report and analytic code) may be available upon request at https://vivili.org/. In cases where clinical trial data and supporting documents are provided pursuant to company policies and procedures, Daiichi Sankyo will continue to protect the privacy of clinical trial participants. Details on data sharing criteria and the procedure for requesting access can be found at this web address: https://vivili.org/ourmember/daiichi-sankyo/

Code availability Not applicable.

Authors’ contributions Nobuyuki Sakai contributed to study design and preparation of manuscript. Masataka Takeuchi, Hirotsoshi Imamura, Norihito Shimamura, Shinichi Yoshimura, Hiroichi Naito, Naoto Kimura, Osamu Masuo, Nobuyuki Hirotsune, Kenichi Morita, Kazu- nori Toyoda, Hiroshi Yamagami and Hideyuki Iishihara contributed to patient enrollment. Shigeru Fujimoto contributed to data analysis and determining outcomes. Takafumi Nakatsu, Naoki Miyoshi and Miharu Suda contributed to study design, statistical analysis and preparation of the manuscript. All authors read and approved the drafts, including the final version for submission, and agree to be accountable for all aspects of the work.
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