Post-marketing surveillance data of thrombomodulin alfa: sub-analysis in patients with sepsis-induced disseminated intravascular coagulation

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Abstract

Background: Thrombomodulin alfa (TM-α, recombinant thrombomodulin) significantly improved disseminated intravascular coagulation (DIC) when compared with heparin therapy in a phase III study. Post-marketing surveillance of TM-α was performed to evaluate the effects and safety in patients with sepsis-induced DIC.

Methods: From May 2008 to April 2010, a total of 1,787 patients with sepsis-induced DIC treated with TM-α were registered. DIC was diagnosed based on the Japanese Association for Acute Medicine (JAAM) criteria. The DIC resolution and survival rates on day 28 after the last TM-α administration, and changes in DIC, systemic inflammatory response syndrome (SIRS), and sequential organ failure assessment (SOFA) scores and coagulation and inflammation markers were evaluated.

Results: The most frequent underlying disease was infectious focus-unknown sepsis (29.8%). The mean ± SD values of age, dose, and the duration of TM-α administration were 64.7 ± 20.3 years, 297.3 ± 111.4 U/kg/day, and 5.6 ± 3.4 days, respectively. A total of 1,320 subjects (73.9%) received combined administration with other anticoagulants. Both coagulation and inflammation markers, such as fibrin/fibrinogen degradation products, prothrombin time ratio, thrombin-antithrombin complex, and C-reactive protein, as well as JAAM DIC, SIRS, and SOFA scores, significantly and simultaneously decreased after TM-α administration (p < 0.001). DIC resolution and 28-day survival rates were 44.4% and 66.0%, respectively. The 28-day survival rate decreased significantly according to the duration of DIC before TM-α administration (p < 0.001). Total adverse drug reactions (ADRs), bleeding ADRs, and serious bleeding adverse events occurred in 126 (7.1%), 98 (5.5%), and 121 (6.8%) subjects, respectively. On day 28, after the last TM-α administration available for an antibody test, only one patient was positive for anti-TM-α antibodies (0.11%).

Conclusion: Our results suggest that TM-α is most effective for treating patients with sepsis-induced DIC when administered within the first 3 days after diagnosis.

Keywords: Anticoagulant, JAAM criteria, Sepsis, SIRS, SOFA score

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Background
In severe sepsis, disseminated intravascular coagulation (DIC) occurs in about 35% of patients [1]. The mortality rate is higher in patients with DIC (40% to 46.2%) than in patients without DIC (22.2% to 26.5%) [2,3]; thus, the prognosis of sepsis is considered to be affected by the presence of DIC.

Thrombomodulin (TM) is a thrombin receptor on the endothelial cell surface that plays an important role in the regulation of intravascular coagulation [4]. Thrombin binds to TM, and the thrombin-TM complex then activates protein C to form activated protein C (APC), which cleaves and inactivates factors Va and VIIIa in the presence of protein S. Therefore, TM acts as a negative feedback regulator according to the amount of excessively generated thrombin. In addition to its anticoagulant properties, TM has an anti-inflammatory effect through the generation of APC, and more, directly regulates high mobility group box-1 (HMGB-1) [5] and endotoxin [6] through the lectin-like domain, independent of protein C activation.

Thrombomodulin alfa (TM-α; Recomodulin® Injection, Asahi Kasei Pharma Corporation, Tokyo, Japan) is a recombinant, human, soluble TM possessing an extracellular domain that includes a functional domain. The effects of TM-α on DIC have been examined in a multicenter, randomized, clinical trial in Japan [7,8]. In patients with DIC associated with infection or hematological malignancy, resolution rates for DIC and bleeding symptoms were significantly better in the TM-α group than in the heparin group. TM-α was approved for use as a curative medicine for DIC for the first time in Japan in 2008. Since the launch of this product, a clinical use survey was conducted by continuous registration of all cases using TM-α since the launch of this product, a clinical use survey was conducted by continuous registration of all cases using TM-α. TM-α can be administered once a day at a dose of 380-U/kg body weight [7]. Since TM-α is excreted by the kidney, the dose of TM-α administered to subjects with renal dysfunction was reduced to 130-U/kg body weight. TM-α was infused after DIC was diagnosed, but the start of TM-α administration was not defined; therefore, each physician was free to determine the initial timing of TM-α infusion. In this PMS study, no limitation was placed on the period of TM-α administration and each physician individually judged the completion of TM-α administration. There was no limitation on the use of other anticoagulants, including antithrombin substitution, synthetic protease inhibitors (gabexate mesylate or nafamostat mesylate), heparin derivatives (heparin, dalteparin, or danaparoid sodium), or blood preparations such as fresh frozen plasma (FFP) and platelet concentrate, before and/or after TM-α administration.

Serum samples collected at baseline and on day 28 after the last TM-α administration were tested by enzyme-linked immunosorbent assay (ELISA) for anti-TM-α antibodies. This PMS study was conducted in accordance with the guidelines for Good Post-Marketing Surveillance Practices as required by the Japanese Ministry of Health, Labor, and Welfare. All subjects were treated according to the attending physicians’ decisions, and no limitations were placed on the concomitant use of other anticoagulants or medicine for the treatment of underlying diseases and complications. In addition, personal data anonymization was carried out upon data collection. Therefore, approval of this surveillance by ethical committees and institutional review boards or informed consent acquisition was not necessary.

Methods
Study design and patient population
The subjects in the present observational study were a subset of those in the PMS reported by Mimuro et al. [9]. In response to requests for methods to improve the early diagnosis of DIC accompanying infections in the fields of emergency and ICU care, the JAAM DIC criteria aimed at achieving early diagnosis were announced in 2006 by the DIC Special Committees [10]. A total of 1,787 sepsis-induced DIC subjects were analyzed (Figure 1). From May 2008 to April 2010, 1,787 patients were registered in this study from 364 institutes, and DIC was diagnosed based on a score of 4 points on the JAAM DIC scoring system (see Appendix) [10]. The exclusion criteria were as follows: patients without SIRS (SIRS score ≤ 1), missing data for complete analysis, those with an unknown outcome, intracranial, respiratory, or continuous gastrointestinal bleeding, hypersensitivity to this agent, and pregnancy.

Data collection
Organ dysfunction was assessed using the SOFA score. SIRS, sepsis, and septic shock were defined according to the American College of Chest Physicians/Society of Critical Care Medicine consensus conference [11] and Surviving Sepsis Campaign Guidelines 2008 [12], respectively. DIC resolution was defined as a total JAAM DIC score ≤ 3 by the day after the last TM-α administration. The JAAM DIC resolution rate was calculated using data from 1,152 subjects who were categorized as either
‘resolution’ or ‘non-resolution’. The survival rate was calculated based on the number of subjects who were alive 28 days after the initial TM-α administration (n = 1,771). Subjects were followed until day 28 after the last administration. At the start of TM-α administration, information about the following 14 baseline characteristics was collected: age, sex, severity of underlying disease, history of bleeding, duration of DIC, kidney injury, liver injury, bleeding or symptoms of organ dysfunction, platelet count, fibrin/fibrinogen degradation products (FDP) or fibrinogen or antithrombin levels, and PT ratio. Furthermore, the JAAM DIC, SIRS, and SOFA scores were added as a factor relevant to sepsis [10]. Multiple logistic-regression analysis was then conducted using a total of 15 baseline characteristics excluding ‘severity of underlying disease’ and ‘liver injury’, which are subjective observations.

Blood collection for clinical laboratory tests and coagulation studies was performed at the following three time points: before, during, and after TM-α administration. However, the TM-α administration period was excluded from the data analysis because the time varied by case. Safety data were coded using the preferred terms from version 13.1 of the Japanese version of the Medical Dictionary for Regulatory Activities (MedDRA/J) [13]. Definitions of adverse events (AEs) and adverse drug reactions (ADRs) were based on the International Conference on Harmonization of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) guidelines [14]. The safety evaluation included serious bleeding AEs and all ADRs observed until 28 days after the last TM-α administration.

**Statistical analysis**

In the descriptive analysis of baseline characteristics, numerical data are expressed as means ± standard deviation (SD) or medians (Q1–Q3; interquartile range). Statistical analysis was performed to compare values using the chi-square test, the Wilcoxon signed rank test, and the Cochran-Armitage test. To identify risk factors, DIC resolution and survival at 28 days after starting TM-α administration were stratified and tabulated by the 15 baseline characteristics. Differences in frequency by strata were tested for significance by the chi-square test. Relative risk factors for the DIC resolution rate and the survival rate after TM-α administration for DIC treatment were analyzed by multiple logistic regression analysis. Relative risk factors for serious bleeding AEs were identified using multiple logistic regression analysis. Odds ratios (ORs) with

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*Figure 1 Subject disposition. DIC disseminated intravascular coagulation, JAAM Japanese Association for Acute Medicine, SIRS systemic inflammatory response syndrome, TM-α thrombomodulin alfa.*
95% confidence intervals (CIs) for selected variables were calculated. A value of $p < 0.05$ was considered significant. Statistical testing was performed with JMP version 9.0 software (SAS Institute, Cary, NC, USA).

### Results

#### Subject demographics

A total of 1,787 patients were enrolled. Subject demographic characteristics are shown in Table 1. The mean ±

| Item                                      | Subjects (n) | Rate (%) |
|-------------------------------------------|--------------|----------|
| Sex                                       | Male         | 1038     | 58.1    |
|                                           | Female       | 749      | 41.9    |
| Age (years)                               |              |          |         |
| 0–14                                      | 86           | 4.8      |
| 15–64                                     | 559          | 31.3     |
| 65–74                                     | 491          | 27.5     |
| 75–84                                     | 498          | 27.9     |
| ≥ 85                                      | 149          | 8.3      |
| Unknown                                   | 4            | 0.2      |
| Source of sepsis                          |              |          |         |
| Lungs                                     | 346          | 19.4     |
| Abdomen                                   | 396          | 22.2     |
| Urine                                     | 180          | 10.1     |
| Skin                                      | 48           | 2.7      |
| Focus-unknown                             | 533          | 29.8     |
| Others                                    | 284          | 15.9     |
| DIC score (JAAM criteria)                 |              |          |         |
| 4                                         | 216          | 15.4     |
| 5                                         | 368          | 26.3     |
| 6                                         | 325          | 23.2     |
| 7                                         | 191          | 13.6     |
| 8                                         | 300          | 21.4     |
| SOFA score                                |              |          |         |
| 0–7                                       | 266          | 24.4     |
| 8–10                                      | 270          | 24.8     |
| 11–13                                     | 275          | 25.3     |
| ≥ 14                                      | 278          | 25.5     |
| SIRS score                                |              |          |         |
| 2                                         | 394          | 24.4     |
| 3                                         | 708          | 43.8     |
| 4                                         | 513          | 31.8     |
| Other anticoagulant treatment for DIC     | 1,320/1,020  | 73.9/57.1|
| treatment for DIC (Overall/after TM-α administration) | | |
| AT concentrate                            | 876/476      | 49.0/26.6|
| Synthetic protease inhibitors             | 816/333      | 45.7/18.6|
| Heparin derivatives                       | 427/212      | 23.9/11.9|
| Duration (days) of DIC before TM-α        | 1,134        | 68.3     |
| administration                            | 1            | 6.5      |
| 2                                         | 345          | 19.3     |
| 3                                         | 110          | 6.2      |
| 4–6                                       | 59           | 3.3      |
| ≥ 7                                       | 78           | 4.4      |

**Table 1 Subject baseline demographics and characteristics**

AT antithrombin, DIC disseminated intravascular coagulation, JAAM Japanese Association for Acute Medicine, SIRS systemic inflammatory response syndrome, SOFA Sequential Organ Failure Assessment, TM-α thrombomodulin alfa.

#### Figure 2

Changes of coagulation and inflammation markers before and after TM-α administration. $p$ values: Wilcoxon signed-rank test. CRP C-reactive protein, FDP fibrin/fibrinogen degradation products, PLT platelet, PT prothrombin time, TAT thrombin-antithrombin, TM-α thrombomodulin alfa, WBC white blood cell.
SD age was 64.7 ± 20.3 years, and the median age was 70 (59–78, Q1-Q3) years. A total of 647 subjects (36.3%) were over 75 years old. The most frequent underlying disease at the time when TM-α was started was infectious focus-unknown sepsis (n = 533, 29.8%). The DIC score (median, Q1–Q3) was 6.0 (5.0–7.0). There were 816 (58.3%) subjects with a DIC score over 6 points and thus considered to be critically ill [15]. The SOFA and SIRS scores (median, Q1–Q3) were 11 (8.0–14.0) and 3.0 (3.0–4.0), respectively. Using the SOFA score, 553 subjects (50.8%) were severe cases, scoring over 11 points [16]. In regards to the timing of TM-α administration, 63.5% of subjects were treated with TM-α on the same day that they were diagnosed as having DIC, but 25.5% and 11.1% were started 1 to 2 days and over 3 days after being diagnosed as having DIC, respectively. The mean ± SD duration of TM-α administration was 5.6 ± 3.4 days, and the most frequent duration was 6 days (n = 589). The mean ± SD dose was 297.3 ± 111.4 U/kg/day, whereas the most common doses were near 380 (n = 1,008, 56.5%) and 130 (n = 424, 23.8%) U/kg/day.

Changes in coagulation and inflammation markers

Figure 2 shows the coagulation and inflammation markers before and after TM-α administration. All values are shown as median (Q1–Q3). The median platelet count increased significantly from 5.0 (2.8–7.3) to 7.0 (3.4–12.8) × 10^4/μL (p < 0.001), and plasma levels of FDP (μg/mL), PT ratio, and TAT (ng/mL) decreased significantly from 29.5 (14.6–60.5) to 13.0 (8.0–22.3) (p < 0.001), 1.3 (1.2–1.6) to 1.2 (1.1–1.4) (p < 0.001), and 13.4 (7.0–23.1) to 5.3 (3.2–10.6) (p < 0.001), respectively. In the 1,076 subjects who had not been given platelet concentrates and FFP, the median platelet count increased significantly from 5.6 (3.3–7.7) to 8.5 (4.2–14.5) × 10^4/μL (p < 0.001), and in 966 subjects, the PT ratio decreased from 1.3 (1.2–1.5) to 1.2 (1.1–1.3) (p < 0.001). CRP (mg/dL) levels decreased significantly from 16.7 (9.8–24.0) to 7.7 (3.7–13.2) (p < 0.001), according to the improvement of these coagulation parameters. White blood cell (WBC) counts did not change, even when the subjects who had both WBC and CRP levels measured (n = 1,574) were examined. Fibrinogen (mg/dL) levels decreased slightly, but significantly, from 370 (247–504) to 347 (239–472) (p < 0.001) before and after TM-α administration.

Changes in the DIC, SIRS, and SOFA scores

Changes in the DIC, SIRS, and SOFA scores before and after TM-α administration are shown in Figure 3. The DIC, SIRS, and SOFA scores decreased significantly from 6.0 (5.0–7.0) to 4.0 (2.0–5.0) (p < 0.001), 3.0 (3.0–4.0) to 2.0 (1.0–3.0) (p < 0.001), and 11.0 (8.0–14.0) to 7.0 (4.0–12.0) (p < 0.001), respectively. These findings indicate that recovery of the coagulation disorders and attenuation of both inflammation and organ dysfunction occurred simultaneously during TM-α administration.

DIC resolution rate

Of the 1,152 subjects, DIC resolved in 512 after TM-α administration; therefore, the DIC resolution rate was 44.4%. The resolution rates decreased significantly in proportion to the severity of the DIC and SOFA scores before TM-α administration (p < 0.001) (Table 2). Comparing the subjects who were injected with 130 and 380 U/kg of TM-α, the resolution rates were 29.8% (78/262) and 51.7% (341/660), respectively; this difference was significant (p < 0.001). Other anticoagulants before TM-α administration did not affect the resolution rate (present 43.5% (225/517) vs. absent 45.2% (287/635), p = 0.724). A multiple logistic regression analysis showed that

![Figure 3 Changes in JAAM DIC, SIRS, and SOFA scores before and after TM-α administration. p values: Wilcoxon signed-rank test. DIC disseminated intravascular coagulation, JAAM Japanese Association for Acute Medicine, SIRS systemic inflammatory response syndrome, SOFA Sequential Organ Failure Assessment.](http://www.jintensivecare.com/content/2/1/30)
fibrinogen, JAAM DIC, and SOFA scores before TM-α administration were significant independent factors affecting the DIC resolution rate (Table 3).

Survival rate

With respect to overall mortality, 1,144 of the 1,771 subjects survived 28 days after TM-α administration; therefore, the overall mortality rate was 35.4%. The survival rate decreased significantly in proportion to the duration of DIC before TM-α administration (p < 0.001) (Figure 4). More precisely, the 28-day survival rate was 66.4% when TM-α was injected on the same day that DIC was diagnosed, whereas it was 48.5% when TM-α was started 4 days later, and 31.1% when it was started 7 or more days after DIC was diagnosed. These differences were significant (p = 0.033 and p < 0.001, respectively). The survival rate decreased significantly in proportion to the severity of the SOFA score before TM-α, but not the DIC score (Table 4). The 28-day survival rate was 88.4% (449/508) in subjects whose DIC diagnosed by the JAAM criteria resolved after administration of TM-α, whereas it was 54.1% (343/643) in the subjects whose DIC continued. Therefore, resolution of DIC diagnosed by the JAAM criteria after TM-α administration strongly reflects the 28-day survival rate. Use of other anticoagulants before and after TM-α administration did not affect the survival rate (present vs. absent: 63.8% (485/760) vs. 65.2% (659/1011), p = 0.782, and 62.5% (288/461) vs. 65.3% (856/1310), p = 0.606, respectively). Survival rates in the subjects treated with 130 and 380 U/kg of TM-α were 58.5% (245/419) and 68.8% (688/1,000), respectively, with a tendency for improvement with 380 U/kg (p = 0.085). A multiple logistic regression analysis revealed that sex, duration of DIC, fibrinogen, and SOFA score before TM-α administration were significant independent factors related to mortality (Table 5).

### Table 2 The JAAM DIC resolution rate of DIC and SOFA scores before TM-α administration

| Factor | Total subjects (n) | DIC resolution | p value |
|--------|-------------------|----------------|---------|
|        | Subjects (n) Rate (%) |
| JAAM DIC score before TM-α administration | Total 1,089 | 488 | 44.8 | - |
| 4 | 164 | 93 | 56.7 | < 0.001 |
| 5 | 285 | 146 | 51.2 |  |
| 6 | 259 | 110 | 42.5 |  |
| 7 | 153 | 60 | 39.2 |  |
| 8 | 228 | 79 | 34.6 |  |
| SOFA score before TM-α administration | Total 768 | 348 | 45.3 | - |
| 0–7 | 173 | 104 | 60.1 | < 0.001 |
| 8–10 | 192 | 94 | 49.0 |  |
| 11–13 | 209 | 91 | 43.5 |  |
| ≥ 14 | 194 | 59 | 30.4 |  |

*Chi-square test.

### Table 3 Multiple logistic regression analysis of risk factors affecting the JAAM DIC resolution rate

| Item | Odds ratio (95% CI) | p value |
|------|---------------------|---------|
| Fibrinogen before TM-α administration | 1.002 (1.001–1.003) | < 0.0001 |
| JAAM DIC score before TM-α administration | 0.836 (0.743–0.939) | 0.0026 |
| SOFA score before TM-α administration | 0.894 (0.857–0.932) | < 0.0001 |

A total of 693 DIC patients diagnosed by JAAM DIC criteria were analyzed by multiple logistic regression analysis; CI confidence interval, DIC disseminated intravascular coagulation, JAAM Japanese Association for Acute Medicine, SOFA Sequential Organ Failure Assessment, TM-α thrombomodulin alfa.

### Table 4 The 28-day survival rate after TM-α administration of DIC and SOFA scores before TM-α administration

| Factor | Total subjects (n) | Survival rate | p value |
|--------|-------------------|---------------|---------|
| JAAM DIC score before TM-α administration | Total 1,387 | 915 | 66.0 | - |
| 4 | 215 | 147 | 68.4 | 0.630 |
| 5 | 363 | 248 | 68.3 |  |
| 6 | 323 | 208 | 64.4 |  |
| 7 | 191 | 121 | 63.4 |  |
| 8 | 295 | 191 | 64.7 |  |
| SOFA score before TM-α administration | Total 1,077 | 712 | 66.1 | - |
| 0–7 | 263 | 214 | 81.4 | < 0.001 |
| 8–10 | 266 | 192 | 72.2 |  |
| 11–13 | 270 | 174 | 64.4 |  |
| ≥ 14 | 278 | 132 | 47.5 |  |

*Chi-square test.
Safety
ADRs were observed in 126 subjects (7.1%), and bleeding ADRs and serious bleeding AEs occurred in 98 (5.5%) and 121 (6.8%) subjects, respectively (Table 6). The frequencies of bleeding ADRs and serious bleeding AEs in subjects treated with 130 and 380 U/kg were 5.9% and 4.9%, and 9.4% and 5.2%, respectively. Therefore, it is possible that the occurrence of these ADRs and AEs did not depend on the TM-α dose; additional research is needed to clarify this point. Of the 1,787 subjects, 121 had serious bleeding AEs, with gastrointestinal bleeding being the most frequently observed serious bleeding AE (2.7%, \( n = 48 \)), followed by respiratory (1.0%, \( n = 18 \)) and intracranial (0.8%, \( n = 15 \)) bleeding, wound bleeding (0.6%, \( n = 11 \)), and injection site bleeding (0.5%, \( n = 9 \)).

To investigate the risk factors before TM-α administration for serious bleeding AEs, a multiple logistic regression analysis was performed (Table 7). History of bleeding risk and pre-existing bleeding symptoms were especially significant independent risk factors for serious bleeding AEs (\( p < 0.0001 \), respectively). Of the 1,787 subjects in this study, 285 (15.9%) and 266 (14.9%) had a history of bleeding risk and pre-existing bleeding symptoms before TM-α administration, respectively. Therefore, in the 1,282 subjects with neither a history of bleeding risk nor pre-existing bleeding symptoms, the frequencies of bleeding ADRs and serious bleeding AEs were 3.7% (\( n = 48 \)) and 4.3% (\( n = 55 \)), respectively. Although the incidence of bleeding is relatively small, it is not rare. Bleeding ADRs did not affect survival [present vs. absent: 62.2% (61/98) vs. 64.7% (1,083/1,673), \( p = 0.815 \)], but serious bleeding AEs did [present vs. absent: 38.0% (46/121) vs. 66.5% (1,098/1,650), \( p = 0.001 \)]. No subject was found to have antibodies that neutralized TM-α activity.

**Discussion**
In this PMS sub-population analysis, in which DIC was diagnosed according to the JAAM DIC criteria and TM-α was administered at 4 days or later, the 28-day survival rate was significantly decreased as compared with TM-α administration within 3 days of diagnosis. These findings suggest that early use of TM-α could be useful for treating sepsis-induced DIC.

The survival rate in this PMS study was somewhat lower than that of the phase III clinical trial [7]. In the phase III clinical trial, patients on dialysis or with renal dysfunction were excluded because of severely impaired drug excretion from the kidney. In the present PMS sub-population analysis, subjects with renal dysfunction had their dose reduced to 130 U/kg according to the approved prescribing information, because TM-α is predominantly excreted by the kidney. The DIC resolution and survival rates were significantly lower in subjects given 130 U/kg (29.8% and 58.5%, respectively) than in those given 380 U/kg (51.7% and 68.8%, respectively) TM-α. However, those given the lower dose had renal dysfunction and were, therefore, more severely ill than those given the higher dose. Indeed, the subjects’ SOFA scores before TM-α administration were 12.4 ± 4.0 with the lower dose and 9.7 ± 3.9 with the higher dose. This may be the one reason why the subjects treated with the lower dose had a poor DIC resolution rate. As a result, it appears that as the DIC period extended, subjects treated with the lower dose had a higher mortality.

The interaction between inflammation and coagulation involves significant cross-talk between systems [17], and the anti-inflammatory and anticoagulant effects of TM-α have been reported by Ito et al. [18], Yamakawa et al. [19] reported improvement of rapid CRP and FDP with TM-α.
Table 8 JAAM DIC criteria

| Systemic inflammatory response syndrome criteria* | Score |
|--------------------------------------------------|-------|
| ≥ 3                                             | 1     |
| 0–2                                             | 0     |
| Platelet count, ×10^9/μL                         |       |
| <8 or >50% decrease within 24 h                  | 3     |
| ≥8 and <12; or >30% decrease within 24 h         | 1     |
| ≥12                                             | 0     |
| Prothrombin time (value of subject/normal value) |       |
| ≥1.2                                             | 1     |
| <1.2                                             | 0     |
| Fibrin/fibrinogen degradation products, μg/mL    |       |
| ≥25                                             | 3     |
| ≥10 and <25                                     | 1     |
| <10                                             | 0     |
| Diagnosis                                        |       |
| ≥4 points                                       | DIC   |

Scoring system for disseminated intravascular coagulation (DIC) established by the Japanese Association for Acute Medicine. *Criteria for systemic inflammatory response syndrome: temperature > 38°C or < 36°C; heart rate > 90 beats/min; respiratory rate > 20 breaths/min or PaCO2 < 32 torr (< 4.3 kPa); white blood cell counts > 12,000 cells/μL, < 4,000 cells/μL, or 10% immature (band) forms.

as compared with a non-treated TM-α control group. Recently, TM-α improved the elevated circulating levels of HMGB-1 in patients with sepsis-induced DIC. The rate of change in HMGB-1 was 48.1% in the TM-α-treated group and 213% in the non-TM-α-treated group (p = 0.086) [20]. Taken together, the anti-inflammatory effect of TM-α may explain the attenuation of HMGB-1 followed by reduction of CRP. These mechanisms, in part, suggest that TM-α improves outcomes in patients with sepsis-induced DIC. Indeed, Yamakawa et al. reported that the 90-day survival rate was improved by TM-α administration in these DIC patients [21].

On the other hand, as for the present result, the WBC count in association with inflammation was not affected by TM-α administration. WBC counts in patients with severe infection sometimes show levels below the normal range, which then increase according to the attenuation of inflammation. These phenomena may, in part, explain the unchanged WBC levels before and after TM-α administration.

The rates of serious bleeding events and intracranial hemorrhage were somewhat high in the present PMS sub-population analysis. Khan et al. reported that both serious bleeding events and intracranial hemorrhage in patients receiving activated protein C were significantly higher in the retrospective observation studies than in the randomized controlled trial (RCT) [22]. A recent TM-α global phase IIB study of sepsis-induced DIC reported that there were no differences between TM-α and placebo groups in the prevalence of serious major bleeding events (5.1% in the TM-α group vs. 4.6% in the placebo group) [23]. In the present PMS sub-population analysis, bleeding ADRs did not affect survival, but serious bleeding AEs did. Serious bleeding may partially depend on the severity of DIC, because the fibrinogen level before TM-α administration affected bleeding as a serious AE on multiple logistic regression analysis.

In order to estimate the frequency of anti-TM-α antibody production, especially neutralizing antibody that neutralizes the activity of TM-α and/or endothelial cell thrombomodulin, anti-TM-α antibody was measured before, and on day 28 after the last TM-α administration by ELISA for anti-TM-α antibodies. In the 1,787 subjects, there were 945 serum samples taken on day 28 after the last TM-α administration available for an antibody test, and only one patient was positive for anti-TM-α antibodies (0.11%). However, the subject was not found to possess antibodies that neutralized TM-α activity in an in vitro study. Thus, it appears that treatment with TM-α in this subject did not decrease the effect of TM-α.

There were several limitations in the present PMS sub-population analysis. First, this PMS study was an examination of a single arm with no comparison arms. Second, this PMS study was performed under daily clinical practice conditions, with restrictions on neither the treatment of underlying diseases nor the usage of other anticoagulants. However, this investigation offers new information about the effect and safety profile of TM-α for sepsis-induced DIC in daily clinical practice, and it collected prospective data and analyzed sepsis-induced DIC diagnosed according to the JAAM DIC criteria, which was not fully elucidated by the phase III clinical study. The findings of a recent retrospective cohort study revealed that TM-α administration improved sepsis-induced DIC diagnosed according to the JAAM DIC criteria [24]. Despite these limitations, our findings may have important implications. Further study is needed to confirm these findings.

Conclusions
The present findings suggest that TM-α is most effective for treating patients with sepsis-induced DIC when administered within the first 3 days after diagnosis. To verify the curative effect of TM-α on sepsis-induced DIC, a large-scale RCT is required.

Appendix
The JAAM DIC criteria are presented in Table 8.

Abbreviations
AE: adverse event; ADR: adverse drug reaction; APC: activated protein C; AT: antithrombin; CI: confidence interval; CRP: C-reactive protein; DIC: disseminated intravascular coagulation; ELISA: enzyme-linked
immunosorbent assay; FDP: fibrin/fibrinogen degradation products; FFP: fresh frozen plasma; HMGB-1: high mobility group box-1; ICU: intensive care unit; JAAM: Japanese Association for Acute Medicine; MedDRA/J: Medical Dictionary for Regulatory Activities; OR: odds ratio; PLT: platelet; PMS: post-marketing surveillance; PT: prothrombin time; RCT: randomized controlled trial; SD: standard deviation; SIRS: systemic inflammatory response syndrome; SOFA: Sequential Organ Failure Assesment; TAT: thrombin-antithrombin; TM: thrombomodulin; TM-α: thrombomodulin alfa; WBC: white blood cell.

Competing interests
This surveillance study was supported by the Japanese Society on Thrombosis and Hemostasis. O.N. and R.T. are employees of Asahi Kasei Pharma Corporation. S.G. consulted for Asahi Kasei Pharma America (ART 123 Program Advisory Board member). H.I. received payment for educational presentations from Asahi Kasei Pharma. The other authors declare that they have no competing interests.

Authors' contributions
YE, HI, DS, RT, and SG conceived and designed the research; DS and ON analyzed the data; YE wrote the manuscript; JM, H Takahashi, H Tsuchi, TM, YE, IK, and YS were involved in drafting the manuscript and revising it critically for important intellectual content; and YS provided important intellectual content and gave final approval. All authors contributed to the revision of the manuscript. All authors read and approved the final manuscript.

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