Phase II trial of aflibercept with FOLFIRI as a second-line treatment for Japanese patients with metastatic colorectal cancer

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Aflibercept targets vascular endothelial growth factor. The present study involved assessing the efficacy, safety and pharmacokinetics of aflibercept plus 5-fluorouracil/levofolinate/irinotecan (FOLFIRI) as a second-line treatment for metastatic colorectal cancer (mCRC) in Japanese patients. Aflibercept (4 mg/kg) plus FOLFIRI was administered every 2 weeks in 62 patients with mCRC until disease progression, unacceptable toxicity or patient withdrawal. Tumors were imaged every 6 weeks. The primary endpoint was objective
1 | INTRODUCTION

Colorectal cancer is the third most commonly occurring cancer worldwide and the second leading cause of cancer-related deaths. In Japan, it is the most commonly occurring cancer in both sexes combined. The 5-year relative survival rate for metastatic colorectal cancer (mCRC) is approximately 12%-13%. Thus, it is important to develop new, more effective therapies.

Vascular endothelial growth factor (VEGF) is overexpressed in primary colon tumors that have metastasized compared to those that have not. Furthermore, high VEGF expression predicts poor relapse-free and overall survival of individuals with colorectal cancer. For this reason, VEGF has in recent years become a target for anti-cancer therapies, in combination with standard chemotherapy regimens folinic acid/5-fluorouracil (5-FU)/oxaliplatin (FOLFOX), FOLFOX/irinotecan (FOLFOXIRI) or 5-FU/leovolinate/irinotecan (FOLFIRI). One anti–VEGF agent, bevacizumab, specifically blocks VEGF-A and, in combination with FOLFOX, FOLFOXIRI, or FOLFIRI, has been shown to increase survival of patients with mCRC.

Aflibercept, also known as VEGF-trap or ziv-aflibercept, is a relatively new anti-VEGF agent. It is a recombinant fusion protein containing portions of the extracellular domains of human VEGF receptors 1 and 2. Unlike bevacizumab, which binds only to VEGF-A, aflibercept binds to VEGF-A, VEGF-B and placental growth factor, thus blocking their downstream activity.

Several clinical trials have demonstrated the relative safety and efficacy of aflibercept plus FOLFIRI. The largest trial to date was an international randomized double-blind phase III study conducted outside of Japan (VELOUR study [NCT00561470]), which consisted of 1226 patients with mCRC who had previously received oxaliplatin. Aflibercept plus FOLFIRI significantly improved both overall survival (OS; 13.50 vs 12.06 months, \( P = .0032 \)) and progression-free survival (PFS; 6.90 vs 4.67 months, \( P = .0001 \)) compared to placebo plus FOLFIRI. Likewise, in a recently published randomized Phase III study of patients from the Asia-Pacific region, aflibercept plus FOLFIRI improved both OS (14.59 vs 11.93 months, hazard ratio: .794) and PFS (6.90 vs 4.67 months, hazard ratio: .629) compared to placebo plus FOLFIRI. In Japanese patients with mCRC, a Phase I dose-escalation study of aflibercept plus FOLFIRI (NCT00921661) was conducted in Japanese patients with mCRC. No dose-limiting toxicities or major safety issues were observed for 4 mg/kg aflibercept, the standard dose.

The objectives of the current phase II study were to assess the efficacy, safety and pharmacokinetics (PK) of aflibercept plus FOLFIRI as a second-line treatment for mCRC in Japanese patients.

2 | METHODS

2.1 | Patients

Inclusion criteria were: histologically or cytologically proven adenocarcinoma of the colon or rectum; measurable disease, per Response Evaluation Criteria in Solid Tumors (RECIST) version 1.1; inoperable metastatic disease; 1 prior chemotherapeutic, oxaliplatin-containing regimen for metastatic disease, during which or within 6 months after completion of which the disease progressed or patients...
relapsed; Eastern Cooperative Oncology Group performance status of 0 or 1; adequate organ function; and signed, dated informed consent. Exclusion criteria included active infectious disease, gastrointestinal ulcer, bleeding, urine protein-creatinine ratio >1, and uridine-5-diphospho-glucuronosyltransferase 1A1 (UGT1A1) genotype of *6/*6, *28/*28 or *6/*28.

2.2 | Study design

This was a prospective, multicenter, open-label, single-arm study (Figure 1). All patients received the following treatment regimen once every 2 weeks: aflibercept (4 mg/kg) over 1-2 hours by intravenous (i.v.) infusion; then levofofinate (200 mg/m²) over 120 ± 20 minutes, plus irinotecan (180 mg/m²) over 90 ± 15 minutes, simultaneously by i.v. infusion; then 5-FU (400 mg/m²) as a bolus over 2-4 minutes; and then 5-FU (2400 mg/m²) over 41-46 hours by continuous i.v. infusion. Treatment was given until disease progression, unacceptable toxicity or patient withdrawal.

The primary endpoint was objective response rate (ORR). Secondary endpoints were PFS, OS, safety and PK.

2.3 | Efficacy assessments

Tumors were imaged every 6 ± 1 weeks and at the end-of-treatment visit 30 ± 3 days after the last study treatment. In the post-treatment follow-up period, tumors were imaged every 6 ± 1 weeks until disease progression. Survival status was determined every 2 months. ORR was defined as the percentage of patients with either a complete response (CR) or partial response (PR) to study treatment, determined based on tumor assessment by an independent radiological review committee (IRRC) using RECIST version 1.1 criteria. PFS was defined as the time interval from the first study treatment administration to either the first observation of radiologically documented disease progression, determined based on tumor assessment by the IRRC, or death due to any cause, whichever came first. OS was defined as the time interval from the first study treatment administration to death due to any cause.

In an exploratory analysis, ORR, PFS and OS were compared in patients with mutated KRAS (exon 2) vs those with wild-type KRAS and in patients with left-sided primary tumors (descending colon, sigmoid colon, rectosigmoid colon and/or rectum) vs those with right-sided primary tumors (caecum, ascending colon and/or transverse colon).
2.4 | Safety assessments

Safety assessments were performed as shown in Figure 1 and included physical examination, evaluation of laboratory data and assessment of adverse events. Laboratory safety tests were performed at baseline, at every visit before treatment administration and at the end-of-treatment visit, and included hematology, biochemistry, urinalysis, coagulation and any other tests as clinically indicated. Laboratory abnormalities were recorded as adverse events only if they led to study treatment discontinuation or modification (eg, dose reduction, cycle delay or omission) and/or were serious (ie, were life-threatening and/or resulted in hospitalization, disability and death). Adverse events assessed included treatment-emergent adverse events (TEAE), serious adverse events and death.

For immunogenicity evaluation, blood samples were collected before aflibercept infusion in treatment cycles 1 and 3, at 30 ± 3 days and 90 ± 7 days after the last aflibercept infusion, and in cases of infusion-related allergic reaction (Grade ≥ 2) or proteinuria (>3.5 g/24 hours or of renal origin associated with hematuria). The presence of anti-aflibercept antibodies was evaluated in serum using a validated non-quantitative titer-based bridging immunoassay. If the result was positive, then the presence of aflibercept-neutralizing antibodies was evaluated using a non-quantitative competitive ligand-binding assay.

2.5 | Population pharmacokinetics

A population PK approach was used to estimate individual PK parameters for free and VEGF-bound aflibercept in all 62 patients. Blood samples were obtained during treatment cycle 1: pre-treatment, before the end of infusion (EOI) of aflibercept (1 hour), and 3, 23 and 335 hours (Day 14) after the EOI of aflibercept. Blood samples were also obtained pre-dose of every odd-numbered cycle, and 30 and 90 days after the last administration of aflibercept.

Plasma concentrations were measured by validated enzyme-linked immunosorbent assays. Concentrations of VEGF-bound
aflibercept were expressed as the free aflibercept equivalent by multiplying by 0.717, the ratio of the molecular weights of free and VEGF-bound aflibercept. The lower limits of quantification (LLOQ) were 15.6 and (adjusted) 31.5 ng/mL, respectively.

The PK parameters were maximum concentration \( (C_{\text{max}}) \), area under the curve over the dosing interval \( (\text{AUC}_{0-14 \text{ day}}) \), total body clearance \( (\text{CL}) \) and volume of distribution at steady state \( (V_{\text{ss}}) \) for free aflibercept, and \( \text{CL} \) for VEGF-bound aflibercept.

### 2.6 Non-compartmental pharmacokinetics

Pharmacokinetics parameters were calculated for irinotecan, its active metabolite SN-38, and 5-FU in the first 10 patients by non-compartmental analysis (PKDMS Version 2 running with WinNonLin Professional, Version 5.2.1, PharSight, Raleigh-Durham, NC, USA). Blood samples for irinotecan and SN-38 were obtained before aflibercept infusion, just before EOI of irinotecan (1.5 hours), and 2, 4.5 and 23 hours after the start of irinotecan infusion during cycle 1. Blood samples for 5-FU were obtained before the start of aflibercept infusion and 2.5, 21 and 45 hours after the start of 5-FU infusion during cycle 1. Concentrations were measured using validated electrospray liquid chromatography tandem mass spectrometry for irinotecan, SN-38 and 5-FU (LLOQ: 10.0, 1.0 and 5.0 ng/mL, respectively).
The PK parameters for irinotecan and SN-38 were \( C_{\text{max}} \), time to reach \( C_{\text{max}}(t_{\text{max}}) \), AUC until last quantifiable time point (AUC_{last}), AUC extrapolated to infinity (AUC), terminal elimination half-life (\( t_{1/2}\)), and metabolic ratio based on molecular weight (\( R_{\text{met}}\)). For irinotecan, CL and \( V_{\text{ss}} \) were also estimated. For 5-FU, steady-state concentration during constant-rate infusion (\( C_{\text{ss}} \)) and clearance at steady state (CL_{ss}) were estimated.

2.7 | Statistical analysis

This study aimed to estimate the ORR in Japanese patients with mCRC at a certain precision. In prior studies of patients with mCRC treated in the second line with FOLFIRI or FOLFIRI plus aflibercept, ORR ranged from approximately 10% to 20%.\(^{10-17}\) If the ORR observed in this study was in the same range, then 60 patients would provide precision (range of 95% CI) from 0.16 to 0.20.

The best objective response was summarized with descriptive statistics. ORR with its associated 95% CI was calculated using normal approximation based on the best objective response judged by the IRRC.

The median PFS, median OS, and their associated probability of survival at each time point and 95% CI were estimated using the Kaplan-Meier method. The time points were every 3 months, for a total of 15 months for PFS and 24 months for OS.

For each laboratory parameter, a patient was considered evaluable if ≥1 measurement was available on treatment. Laboratory toxicities were graded from 1 (least severe) to 5 per National Cancer Institute Common Terminology Criteria for Adverse Events (NCI CTCAE) version 4.03 and were summarized as "all grades" or "Grade ≥3." For patients with multiple occurrences of a particular laboratory parameter during the study treatment period, the maximum (worst) grade was used.

Treatment-emergent adverse events were adverse events reported between the first study treatment infusion and 30 days after the last one. TEAE were summarized with respect to frequency and intensity/severity, as graded by the worst NCI CTCAE version 4.03 criteria. All TEAE were coded using the Medical Dictionary for Regulatory Activities version 18.0.

2.8 | Ethical considerations

The study was conducted in accordance with the principles of the Declaration of Helsinki and in compliance with all international and Japanese laws, regulations and guidelines. All aspects of the study were approved by the independent ethics committee and institutional review board. Patients were fully informed of the study and provided written consent. The study was registered with ClinicalTrials.gov with the identifier NCT01882868.

3 | RESULTS

This study started when the first patient was enrolled in July 2013 and closed on the day of database lock in August 2015. Sixty-two patients from 19 clinical sites in Japan were enrolled. Two patients were not evaluable for ORR: one had neither a target nor a non–target lesion.
and one had only non–target lesions at baseline. The demographics and disease characteristics of the 60 evaluable patients appear in Table 1. Fifty evaluable patients (83.3%) had received prior bevacizumab.

As of the final database lock, all patients had discontinued study treatment (Figure 1). The median number of treatment cycles received per patient was 8.0 (range: 1-31), and the median duration of study treatment exposure was 21.8 weeks (range: 2-73 weeks). The median relative dose intensities were 72% for aflibercept, 65% for irinotecan and 71% for 5-FU.

### 3.1 Efficacy

Five of the 60 evaluable patients had PR and none had CR, resulting in an ORR of 8.3% (95% CI: 1.3%-15.3%). In addition, 43 (71.7%) had
stable disease, for an overall DCR of 80.0% (95% CI: 69.9%-90.1%; see also Figure 2).

The other efficacy endpoints were evaluated for all 62 patients. The median PFS was 5.42 months (95% CI: 4.14-6.70; Figure 3). As of the last tumor assessment, 47 (75.8%) patients had documented disease progression, 11 (17.7%) had died without disease progression and 4 (6.5%) had no disease progression. The median OS was 15.59 months (range: 11.20-19.81 months; Figure 4). As of the final cutoff date, 21 (33.9%) patients had survived. The 2-year survival rate was 28.2% (95% CI: 14.8%-41.6%).

Of the 62 patients, 26 had mutant KRAS, 29 had wild-type KRAS and 7 had unknown KRAS status. Forty-seven patients had a left-sided primary tumor and 15 had a right-sided tumor. The ORR, PFS and OS based on KRAS status and primary tumor location are shown in Table S1 and Figures S1 and S2.

After discontinuing the study treatment, 51 (82.3%) patients received ≥1 further anti–cancer therapy, including 39 (62.9%) who received further biologics/small molecules (cetuximab: 9 [14.5%]; bevacizumab: 9 [14.5%]; regorafenib: 15 [24.2%]; panitumumab: 10 [16.1%]).

### 3.2 Safety

Hematological abnormality occurred in most patients: leukopenia (87.1% of patients), neutropenia (85.5%), anemia (82.3%) and thrombocytopenia (62.9%) (Table 2). Of all clinical laboratory abnormalities assessed, an abnormal creatinine level was the most common, affecting 60 patients (96.8%). Proteinuria occurred in 51 patients (82.3%); of these cases, 22 (43.1%) occurred in treatment cycle 1.

All patients had ≥1 TEAE. Fifty-six patients (90.3%) had Grade 3 or 4 TEAE; the most common TEAE were neutropenia, hypertension, diarrhea and decreased appetite (Table 3). Furthermore, 20 patients had ≥1 serious adverse event. Forty-one (66.1%) patients died due to disease progression, all >30 days after the last study treatment administration. No patients died as a result of treatment.

One patient was positive for anti–aflibercept antibodies at baseline (and positive in the neutralizing antibody assay); however, all subsequent samples from this patient were negative.

### 3.3 Pharmacokinetics

Population PK data (free and VEGF-bound aflibercept) and non–compartmental PK data (irinotecan, SN-38 and 5-FU) appear in Table 4. Plasma concentrations for all 5 analytes are shown in Figure 5.

### 4 DISCUSSION

This study examined the efficacy, safety and PK of aflibercept plus FOLFIRI in Japanese patients with mCRC that was refractory or intolerant to a first-line oxaliplatin-containing regimen. The primary endpoint, ORR by the IRRC, was 8.3% (95% CI: 1.3%-15.3%). The ORR was lower than that in the VELOUR study (19.8%; 95% CI: 16.4%-23.2%). Possible reasons for this discrepancy are differences between the study populations and the limited size of our study. Consistent with a lower ORR, the PFS in the current study was slightly shorter than that in the VELOUR study: 5.42 months (95% CI: 4.14-6.70) vs 6.90 months (6.51-7.20) overall. Based on these findings and the DCR of 80.0% (95% CI: 69.9%-90.1%), adding...
Aflibercept to FOLFIRI yielded results consistent with those of the VELOUR study. The ORR and PFS were also comparable to those in the Asia-Pacific study (ORR = 8.3% [95% CI: 1.3%-15.3%] vs 18% [13.3%-23.5%] and PFS = 5.42 months [4.14-6.70] vs 6.93 months [6.045-7.655]).

The median OS of 15.59 months (95% CI: 11.20-19.81) was similar to the median OS in both the VELOUR study (13.50 months; 95% CI: 12.52-14.95) and the Asia-Pacific study (14.59 months; 95% CI: 13.18-16.46), with overlapping 95% CI. Neither the KRAS oncogene nor primary tumor sidedness significantly affected ORR, PFS or OS.

The safety profile was as expected of an anti–VEGF agent and was comparable to that observed in the VELOUR study. Together, the TEAE represent an enhancement of the toxicity profile associated with usage of FOLFIRI. The absence of a sample positive for anti–aflibercept antibodies post–treatment suggests that IV administration of 4 mg/kg aflibercept confers no immunogenicity in patients with mCRC.

The PK values of free aflibercept were comparable to those in previous studies for Chinese patients. Adding aflibercept to FOLFIRI did not significantly affect the PK of irinotecan, SN-38 or 5-fluorouracil: the PK values of these FOLFIRI components were comparable between the current study, in which aflibercept was added, and published studies in which it was not. For example, the clearance of irinotecan in the current study was similar to the values in the Gupta et al study and in the Satoh et al study. The AUC for irinotecan and SN-38 overlaps or is slightly high. The clearance of 5-FU in our study had a relatively large coefficient of variance.

| TABLE 3 Most commonly reported treatment-emergent adverse events |
|---------------------------------------------------------------|
| **Primary system organ class**                               | **n (%)** |
| Preferred term                                               | All grades | Grade ≥3 |
| Any class                                                     | 62 (100)   | 56 (90.3) |
| Blood and lymphatic system disorders                         | 46 (74.2)  | 38 (61.3) |
| Neutropenia                                                   | 46 (74.2)  | 38 (61.3) |
| Metabolism and nutrition disorders                           | 47 (75.8)  | 10 (16.1) |
| Decreased appetite                                            | 46 (74.2)  | 8 (12.9)  |
| Nervous system disorders                                      | 18 (29.0)  | 0        |
| Headache                                                      | 7 (11.3)   | 0        |
| Vascular disorders                                           | 33 (53.2)  | 17 (27.4) |
| Hypertension                                                  | 29 (46.8)  | 17 (27.4) |
| Respiratory, thoracic, and mediastinal disorders              | 45 (72.6)  | 2 (3.2)   |
| Epistaxis                                                     | 25 (40.3)  | 0        |
| Dysphonia                                                     | 18 (29.0)  | 0        |
| Cough                                                         | 7 (11.3)   | 0        |
| Hiccups                                                       | 7 (11.3)   | 0        |
| Gastrointestinal disorders                                    | 56 (90.3)  | 18 (29.0) |
| Diarrhea                                                      | 42 (67.7)  | 12 (19.4) |
| Nausea                                                        | 36 (58.1)  | 2 (3.2)   |
| Stomatitis                                                    | 29 (46.8)  | 5 (8.1)   |
| Vomiting                                                      | 17 (27.4)  | 0        |
| Constipation                                                  | 10 (16.1)  | 0        |
| Abdominal pain                                                | 9 (14.5)   | 0        |
| Skin and subcutaneous tissue disorders                       | 44 (71.0)  | 1 (1.6)   |
| Alopecia                                                      | 30 (48.4)  | 0        |
| Palmar-plantar erythrodysesthesia syndrome                    | 8 (12.9)   | 0        |
| Rash                                                          | 7 (11.3)   | 0        |
| Renal and urinary disorders                                   | 19 (30.6)  | 6 (9.7)   |
| Proteinuria                                                   | 19 (30.6)  | 6 (9.7)   |
| General disorders and administration site conditions          | 44 (71.0)  | 4 (6.5)   |
| Fatigue                                                       | 39 (62.9)  | 3 (4.8)   |
| Pyrexia                                                       | 13 (21.0)  | 0        |

| TABLE 4 Summary of PK parameters in treatment cycle 1 |
|------------------------------------------------------|
| **PK parameters**                                   | **Mean ± SD (CV%)** |
| Free aflibercept                                     |                       |
| $C_{\text{max}}, \mu g/mL$                          | 73.2 ± 10.7 (15%)     |
| $\text{AUC}_{0-14 \text{ d}}, \mu g \cdot \text{ day}/mL$ | 247 ± 41 (17%)        |
| Clearance, L/d                                       | 0.805 ± 0.178 (22%)   |
| $V_{\text{ss}}, L$                                   | 6.20 ± 1.11 (18%)     |
| VEGF-bound aflibercept                               |                       |
| Clearance, L/d                                       | 0.162 ± 0.014 (9%)    |
| Irinotecan                                           |                       |
| $C_{\text{max}}, \text{ng/mL}$                      | 2220 ± 528 (24%)      |
| $\text{AUC}, \text{ng} \cdot \text{h}/mL$           | 17 700 ± 6400 (36%)   |
| $t_{1/2z}, h$                                        | 5.19 ± 0.74 (14%)     |
| Clearance, L/h/m²                                    | 11.1 ± 3.2 (28%)      |
| $V_{\text{ss}}, L/m²$                                | 55.7 ± 16.0 (29%)     |
| SN-38                                               |                       |
| $C_{\text{max}}, \text{ng/mL}$                      | 32.2 ± 11.4 (36%)     |
| $\text{AUC}, \text{ng} \cdot \text{h}/mL$           | 341 ± 72 (21%)        |
| $t_{1/2z}, h$                                        | 10.3 ± 3.1 (30%)      |
| 5-FU                                                |                       |
| $CL_{\text{ss}}, L/h/m²$                             | 72.6 ± 40.4 (56%)     |

$\text{AUC}_{0-14 \text{ d}}$ area under the concentration vs time curve 0-14 d post start of infusion; $CL_{\text{ss}}$, clearance at steady state; $C_{\text{max}}$, maximum plasma concentration observed; CV%, coefficient of variation; 5-FU, 5-fluorouracil; PK, pharmacokinetics; SD, standard deviation; $t_{1/2z}$, terminal elimination half-life; $V_{\text{ss}}$, volume of distribution at steady state.

95% CI: 13.18-16.46), with overlapping 95% CI. Neither the KRAS oncogene nor primary tumor sidedness significantly affected ORR, PFS or OS.

The safety profile was as expected of an anti–VEGF agent and was comparable to that observed in the VELOUR study. Together, the TEAE represent an enhancement of the toxicity profile associated with usage of FOLFIRI. The absence of a sample positive for anti–aflibercept antibodies post–treatment suggests that IV administration of 4 mg/kg aflibercept confers no immunogenicity in patients with mCRC.

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FIGURE 5  Plasma concentrations of study drugs over time in treatment cycle 1. Blood samples were collected from 10 patients before treatment cycle 1 and at various time points throughout cycle 1. Plasma concentrations of (A) free and vascular endothelial growth factor (VEGF)-bound aflibercept were determined using validated enzyme-linked immunosorbent assays. Plasma concentrations of (B) irinotecan, SN-38 and (C) 5-fluourouracil (5-FU) were determined using validated electrospray liquid chromatography tandem mass spectrometry.
(56%), consistent with studies showing that 5-FU plasma clearance is widely variable among patients (reviewed by Lee et al). In previous population pharmacokinetic analyses, no clinically relevant drug-drug interactions between aflibercept and irinotecan or fluorouracil were found. Likewise, no significant drug-drug interactions were found in the current study.

A limitation of this study is its small sample size compared to that of the VELOUR study. In addition, unlike the VELOUR study, the current study had no control arm, so ability to compare the ORR results in the 2 studies is limited. Studies containing larger numbers of Japanese patients are needed to corroborate our findings.

In conclusion, adding aflibercept to FOLFIRI was shown to be beneficial and well-tolerated in Japanese patients with mCRC.

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CONFLICT OF INTEREST

TH received honoraria from Taiho, Chugai, Takeda, Yakult and Merck Serono, a consulting fee from NanoCarrier, and research funding from MSD, Ono, Sanofi, Daiichi Sankyo, Sumitomo Dainippon Pharma, Taiho, Teijin and NanoCarrier. NS received honoraria from Chugai and Eli Lilly, and research funding from Chugai, Eli Lilly, Dainippon Sumitomo, Taiho, MSD, Ono, Dai-ichi Sankyo and Sanofi. TU received honoraria from Merck Serono, Taiho, Chugai and Takeda, and research funding from Sanofi. KY received honoraria from Chugai, Takeda, Yakult, Daiichi-Sankyo, Merck Serono, Bristol, Bayer, Eli Lilly and Taiho, and research funding from BMS and Sanofi. HF received research funding from Sanofi. ST received honoraria from Asahikasei, and research funding from Merck Serono, Ono and Sanofi. YK received honoraria from BMS and Sanofi, and research funding from Eli-Lilly, BMS and Sanofi. TE received honoraria from Eli Lilly, and research funding from Boehringer, Daiichi-Sankyo. Dainippon Sumitomo, Eli Lilly, Merck Serono, MSD, Novartis, Ono and Taiho, and a scholarship from Ono. EO received honoraria from Bayer, Chugai, Eli Lilly, Merck Serono, Taiho, Takeda and Yakult. TY received honoraria from Chugai, Eli Lilly, Merck Serono, Sanofi and Taiho, and research funding from Boehringer Ingelheim, Chugai, Dainippon Sumitomo, GlaxoSmithKline, MSD, Novartis and Sanofi. The affiliated medical institutions of physician authors received research study funding from Sanofi. TS, YS, SZL and CB are employees of Sanofi. Because Sanofi is the company that initiated the clinical trial, the company employees were involved in planning the study and analyzing the data.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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