Phase I study of tirabrutinib (ONO-4059/GS-4059) in patients with relapsed or refractory B-cell malignancies in Japan

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We evaluated the safety, efficacy, pharmacokinetics, pharmacodynamics and predictive biomarkers of tirabrutinib, a second-generation, enhanced-selectivity Bruton's tyrosine kinase inhibitor in Japanese patients with relapsed/refractory B-cell non-Hodgkin lymphoma (B-cell NHL) and chronic lymphocytic leukemia (CLL). This was an open-label, multicenter, phase I study. Seventeen patients (male N = 8) with a median age of 70 years were enrolled in 4 dose cohorts (160 mg once daily [N = 3], 320 mg once daily [N = 3], 480 mg once daily [N = 4] and 300 mg twice daily [N = 7]); 4 patients had continued tirabrutinib administration as of 4 January 2018. The maximum tolerated dose was not reached. Pneumonitis (N = 1) was the dose-limiting toxicity for 300 mg twice daily. Common adverse events (AEs) were rash (35.3%) and vomiting (29.4%). Eight patients (47.1%) developed grade ≥3 AEs: neutropenia (23.5%), anemia (11.8%) and leukopenia (11.8%) were frequent. The overall response rate (≥PR) was 76.5% (13/17 patients), including 4 DLBCL patients with no CD79A/B or MYD88 mutations, and 1 CLL patient with a TP53 mutation, providing promising data for future developments. Of 16 patients with measurable lesions during the screening period, 12 showed ≥50% reductions in tumor diameter. In many patients, the tumor size decreased soon after beginning treatment. The maximum serum concentration for tirabrutinib was 611, 1220, 1280 and 886 ng/mL on Day 1 and 484, 971 and 1940, and 961 ng/mL on Day 28 for Cohorts 1-4, respectively. Tirabrutinib pharmacokinetics were linear, with little accumulation following multiple doses. Tirabrutinib was well tolerated and showed promising efficacy for B-cell NHL/CLL.

KEYWORDS
B-cell malignancy, B-cell non-Hodgkin lymphoma, chronic lymphocytic leukemia, safety, tirabrutinib
1 | INTRODUCTION

Bruton’s tyrosine kinase (BTK) has an important role in B-cell signaling, cell proliferation and survival,1 and is, therefore, an attractive target for therapeutic intervention in B-cell malignancies and autoimmune disorders. Ibrutinib, the first-in-class irreversible BTK inhibitor, is approved in the United States for treatment of mantle cell lymphoma (MCL), chronic lymphocytic leukemia (CLL)/small lymphocytic lymphoma (SLL), Waldenström’s macroglobulinemia (WM) and marginal zone lymphomas. In contrast, ibrutinib is approved in Japan only for the treatment of MCL and CLL/SLL.2-4 Although the overall response rate (ORR) with ibrutinib is high, complete response (CR) is not frequently achieved. Ibrutinib can also lead to clinically significant toxicities in some patients, including bleeding and atrial fibrillation. It is likely that these adverse events (AEs) are partly due to the low selectivity of ibrutinib.5 Thus, an unmet need remains for novel, more specific BTK inhibitors that might improve efficacy, with less toxicity and shorter therapy duration. More specific second-generation BTK inhibitors, including acalabrutinib (ACP-196), tirabrutinib (ONO-4059/GS-4059) and BGB-3111, have been developed to overcome the limitations of first-generation BTK inhibitors, such as “off-target” AEs or the development of resistance.

Tirabrutinib (ONO-4059/GS-4059) is a covalent and selective oral inhibitor of BTK, currently under clinical development for the treatment of CLL, B-cell non−Hodgkin lymphoma (B-cell NHL) and rheumatoid arthritis. Tirabrutinib has demonstrated cytostatic properties in vitro (against the activated B-cell-like diffuse large B-cell lymphoma [ABC-DLBCL] cell line, TMD8) and in vivo (TMD8 transplanted mouse tumor model).6 Furthermore, promising efficacy data have been reported, and tolerability was observed up to 480 mg once daily (QD) in B-cell NHL and 600 mg QD in CLL in a European phase I study (ONO-4059POE001, #NCT01659255).7 However, this European phase I trial did not include Asian patients and, therefore, the tolerability, safety, efficacy, pharmacokinetics (PK) and pharmacodynamics (PD) of tirabrutinib in Japanese patients with B-cell NHL and CLL are unknown.

The present open-label, multicenter, non−randomized phase I study aimed to evaluate the maximum tolerated dose (MTD), safety, efficacy, PK, PD and predictive biomarkers of tirabrutinib monotherapy in Japanese patients with relapsed or refractory B-cell NHL and CLL.

2 | PATIENTS AND METHODS

2.1 | Patients

Patients with relapsed/refractory B-cell NHL and CLL were enrolled in this study. The main inclusion criteria were: age ≥20 years; confirmed histopathological diagnosis and documented history of relapsed or refractory B-cell non−Hodgkin lymphoma (DLBCL, MCL, follicular lymphoma [FL], marginal zone lymphoma or WM/lymphoplasmacytic lymphoma [LPL]) and CLL/SLL for which no curative or high-priority therapy exists; Eastern Cooperative Oncology Group performance status of ≤2; bidimensionally measurable disease >1.5 cm in its largest diameter (except for patients with CLL and WM); and life expectancy ≥90 days.

Exclusion criteria included: uncontrolled autoimmune hemolytic anemia (CLL only); pancreatitis; cerebellar disorders; disease significantly affecting gastrointestinal function; history of severe allergic or anaphylactic reactions; known active infection, including positive result for HIV, HBV (patients who are HBs antigen negative are to be excluded if they are positive for HBs antibody or HBC antibody and HBV-DNA has been detected by quantitative analysis) or HCV; significant uncontrolled concomitant diseases; prior use of standard anti−lymphoma/leukemia therapy or radiation therapy within 28 days of receiving the first dose of tirabrutinib; prior tirabrutinib treatment; pregnancy; and abnormal laboratory values (creatinine clearance < 50 mL/min [mL/min/1.73 m²]; aspartate aminotransferase or alanine aminotransferase ≥2.5 times the upper limit of normal; platelet count < 50 × 10^9/L; neutrophils < 1.0 × 10^9/L; hemoglobin < 8.0 g/dL; and total bilirubin ≥1.5 times the upper limit of normal).

Informed consent was obtained from all patients prior to enrollment. The study protocol and all amendments were reviewed by ethics committees at each participating center. The study was conducted according to the protocol, Good Clinical Practice guidelines, applicable local regulations and the Declaration of Helsinki.

2.2 | Study design

Patients were enrolled from January 2015 to February 2016 in 4 dose cohorts of 160 mg QD, 320 mg QD, 480 mg QD and 300 mg BID (Cohorts 1-4). The study design was a 3 + 3 dose escalation scheme where patients were observed for a 28-day period to identify dose-limiting toxicities (DLT), which are defined as drug-related AEs that occurred in the first 28 days after the first dose. Further information on the definition of DLT is described in Data S1.

Tirabrutinib administration was continued until progressive disease, unacceptable AEs or consent withdrawal. PK was assessed on Day 1 and Day 28 in Cycle 1. Criteria for the discontinuation or interruption, resumption and dose modifications included AEs meeting DLT criteria, AEs not related to tirabrutinib, a patient’s wish to discontinue, not meeting inclusion criteria and meeting exclusion criteria. The full criteria for discontinuation or interruption, resumption, and dose modification are described in Data S1.

2.3 | Assessments

2.3.1 | Safety

Adverse events were evaluated using the Common Terminology Criteria for Adverse Events (Ver. 4.0). Laboratory tests, vital signs and electrocardiograms were also assessed using conventional methods.
2.3.2 | Efficacy

The assessment of best clinical response was based on International Workshop Criteria for B-cell NHL patients, WM patients and CLL patients. Drug efficacy evaluation was based on the following: overall response rate (ORR), CR (CR unconfirmed or CR with incomplete blood count recovery), partial response rate (PR or very good PR), progression-free survival, overall survival, duration of response and event-free survival.

2.3.3 | Pharmacokinetics

Blood for the PK analysis was drawn on Days 1 and 28 of Cycle 1 (pre-dose and at 30 minutes, 1, 2, 3, 4, 6, 8 and 12 hours post-dose) and on Days 2, 8 and 15 of Cycle 1 (pre-dose and at 4 hours post-dose). The maximum plasma concentration \( (C_{\text{max}}) \), time to maximum plasma concentration \( (T_{\text{max}}) \), area under the concentration–time curve (AUC) and \( T_{1/2} \) were assessed (Phoenix WinNonlin Ver.6.2.1).

2.3.4 | Pharmacodynamics

Bruton’s tyrosine kinase phosphorylation inhibition in peripheral blood mononuclear cells was assessed using western blotting analysis at Day 1 of Cycle 1, from a blood sample taken pre-dose and at 2 hours post-dose. Primary antibodies included anti-phosphorylated BTK (pBTK) (Abcam, Cambridge, MA, USA), anti-BTK (Santa Cruz Biotechnology, Santa Cruz, CA, USA) and anti-beta actin (Sigma-Aldrich). Additional details are provided in Data S1.

2.3.5 | Biomarker analysis

For chemokine analysis, blood was drawn: during the screening period (14 days to 1 day before start of administration); on Days 1, 2, 8, 15 and 28 of Cycle 1; and on Day 1 (all before administration of the study drug) of Cycles 3, 4, 5 and 6. Blood samples were analyzed at a central testing facility (including CCL-3, CCL-4, CXCL-4, CXCL-7, CXCL-10, CXCL-12, CXCL-13 and BIM).

For CLL and non-germinal center B-cell-like (non-GCB) DLBCL patients, cell-free DNA from blood samples was used for central analysis of genetic predictive markers (including 17p deletion and mutation, 11q, 13q, trisomy 12 and TP53 mutational status in CLL, and CD79A, CD79B, MYD88 and CARD11 mutational status in DLBCL). Cell-free DNA was extracted using a QIAamp DNA Mini Kit (Qiagen, Valencia, CA, USA), and mutations were determined by next-generation sequencing. Determination of DLBCL subtype was performed locally by immunohistochemistry according to the Hans criteria.

2.4 | Statistical analysis

Safety and efficacy analyses were based on all patients enrolled in this study who received at least 1 tirabrutinib dose. Best overall response was used for response analysis. ORR (CR plus PR, including

| TABLE 1 | Baseline demographics and disease characteristics of patients (N = 17) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Characteristic  | 160 mg QD, N = 3 | 320 mg QD, N = 3 | 480 mg QD, N = 4 | 300 mg BID, N = 7 | Total N = 17    |
| Age Median, years (range) | 70.0 (62-70) | 72.0 (68-72) | 74.5 (68-79) | 71.0 (37-80) | 70.0 (37-80) |
| <65 y, n (%) | 1 (33.3) | 0 | 0 | 1 (14.3) | 2 (11.8) |
| ≥65 y, n (%) | 2 (66.7) | 3 (100.0) | 4 (100.0) | 6 (85.7) | 15 (88.2) |
| Sex Male, n (%) | 3 (100.0) | 1 (33.3) | 1 (25.0) | 3 (42.9) | 8 (47.1) |
| Female, n (%) | 0 | 2 (66.7) | 3 (75.0) | 4 (57.1) | 9 (52.9) |
| Histologic subtype, n (%) | | | | | |
| Non-GCB DLBCL | 0 | 0 | 2 (50.0) | 2 (28.6) | 4 (23.5) |
| DBLBC | 0 | 0 | 0 | 1 (14.3) | 1 (5.9) |
| MCL | 1 (33.3) | 2 (66.7) | 1 (25.0) | 0 | 4 (23.5) |
| FL | 0 | 1 (33.3) | 1 (25.0) | 3 (42.9) | 5 (29.4) |
| WM | 1 (33.3) | 0 | 0 | 1 (14.3) | 2 (11.8) |
| CLL | 1 (33.3) | 0 | 0 | 0 | 1 (5.9) |
| Relapsed or refractory after the latest treatment, n (%) | | | | | |
| Relapsed | 0 | 2 (66.7) | 2 (50.0) | 4 (57.1) | 8 (47.1) |
| Refractory | 3 (100.0) | 1 (33.3) | 2 (50.0) | 3 (42.9) | 9 (52.9) |
| Prior therapies, n (%) | | | | | |
| Prior rituximab | 2 (66.7) | 3 (100.0) | 4 (100.0) | 7 (100.0) | 16 (94.1) |

CLL, chronic lymphocytic leukemia; DLBCL, diffuse large B-cell lymphoma; FL, follicular lymphoma; MCL, mantle cell lymphoma; non-GCB DLBCL, non-germinal center B-cell-like diffuse large B-cell lymphoma; WM, Waldenström’s macroglobulinemia.
### Table 2  AEs reported for ≥10% of total population in all cohorts

| AEs                               | 160 mg QD N = 3, n (%) | 320 mg QD N = 3, n (%) | 480 mg QD N = 4, n (%) | 300 mg BID N = 7, n (%) | Total (all cohorts) N = 17, n (%) |
|------------------------------------|-------------------------|-------------------------|-------------------------|--------------------------|----------------------------------|
|                                   | Any grade | Grade ≥3 | Any grade | Grade ≥3 | Any grade | Grade ≥3 | Any grade | Grade ≥3 | Any grade | Grade ≥3 |
| All AEs                           | 3 (100.0) | 1 (33.3) | 3 (100.0) | 2 (66.7) | 4 (100.0) | 2 (50.0) | 7 (100.0) | 3 (42.9) | 17 (100.0) | 8 (47.1) |
| Rash                              | 1 (33.3) | 0       | 2 (66.7) | 0       | 2 (50.0) | 0       | 1 (14.3)  | 0       | 6 (35.3)  | 0       |
| Vomiting                          | 0         | 0       | 1 (33.3) | 0       | 2 (50.0) | 0       | 2 (28.6)  | 0       | 5 (29.4)  | 0       |
| Neutropenia                       | 1 (33.3) | 1 (33.3)| 2 (66.7) | 2 (66.7)| 1 (25.0) | 1 (25.0) | 0         | 0       | 4 (23.5)  | 4 (23.5) |
| Anemia                            | 0         | 0       | 0         | 0       | 2 (50.0) | 1 (25.0) | 1 (14.3)  | 1 (14.3) | 3 (17.6)  | 2 (11.8) |
| Constipation                      | 0         | 0       | 1 (33.3) | 0       | 1 (25.0) | 0       | 1 (14.3)  | 0       | 3 (17.6)  | 0       |
| Diarrhea                          | 0         | 0       | 1 (33.3) | 0       | 0         | 0       | 2 (28.6)  | 0       | 3 (17.6)  | 0       |
| Nausea                            | 0         | 0       | 0         | 0       | 1 (25.0) | 0       | 2 (28.6)  | 0       | 3 (17.6)  | 0       |
| Malaise                           | 2 (66.7) | 0       | 0         | 0       | 1 (25.0) | 0       | 0         | 0       | 3 (17.6)  | 0       |
| Arthralgia                        | 1 (33.3) | 0       | 1 (33.3) | 0       | 0         | 0       | 1 (14.3)  | 0       | 3 (17.6)  | 0       |
| Insomnia                          | 0         | 0       | 0         | 0       | 0         | 0       | 3 (42.9)  | 0       | 3 (17.6)  | 0       |
| Leukopenia                        | 0         | 0       | 2 (66.7) | 2 (66.7)| 0         | 0       | 0         | 0       | 2 (11.8)  | 2 (11.8) |
| Lymphopenia                       | 0         | 0       | 1 (33.3) | 1 (33.3)| 1 (25.0) | 0       | 0         | 0       | 2 (11.8)  | 1 (5.9)  |
| Hypophosphatemia                  | 0         | 0       | 0         | 0       | 2 (50.0) | 1 (25.0) | 0         | 0       | 2 (11.8)  | 1 (5.9)  |
| Pyrexia                           | 0         | 0       | 0         | 0       | 0         | 0       | 2 (28.6)  | 0       | 2 (11.8)  | 0       |
| Cystitis                          | 0         | 0       | 1 (33.3) | 0       | 0         | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Nasopharyngitis                   | 1 (33.3) | 0       | 0         | 0       | 0         | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Pharyngitis                       | 1 (33.3) | 0       | 0         | 0       | 0         | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Aspartate aminotransferase increased | 0        | 0       | 0         | 0       | 1 (25.0) | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Gamma-glutamyltransferase increased | 0        | 0       | 1 (33.3) | 0       | 0         | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Thrombocytopenia                  | 0         | 0       | 1 (33.3) | 0       | 1 (25.0) | 0       | 0         | 0       | 2 (11.8)  | 0       |
| Hypokalemia                       | 0         | 0       | 0         | 0       | 2 (50.0) | 0       | 0         | 0       | 2 (11.8)  | 0       |
| Myalgia                           | 2 (66.7) | 0       | 0         | 0       | 0         | 0       | 0         | 0       | 2 (11.8)  | 0       |
| Dysgeusia                         | 0         | 0       | 0         | 0       | 1 (25.0) | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Headache                          | 1 (33.3) | 0       | 0         | 0       | 0         | 0       | 1 (14.3)  | 0       | 2 (11.8)  | 0       |
| Upper respiratory tract inflammation | 1 (33.3)| 0       | 1 (33.3) | 0       | 0         | 0       | 0         | 0       | 2 (11.8)  | 0       |

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*Neutropenia* includes *neutropenia* and *neutrophil count decrease.* AE, adverse events.
PR with residual lymphocytosis) was calculated for the entire study population, and for each treatment cohort and histology.

3 | RESULTS

3.1 | Patients and treatment

Overall, 17 patients with a median age of 70 years (range, 37-80) were enrolled (N = 3, 3, 4 and 7 patients in cohorts 1-4, respectively), and 4 patients had continued tirabrutinib administration as of 4 January 2018. The histological subtypes were FL (N = 5), MCL (N = 4), non-GCB DLBCL (N = 4), WM (N = 2), DLBCL (N = 1) and CLL (N = 1). The median number of prior therapies was 3 (range, 1-11), with 94.1% (16/17 patients) of patients having had prior exposure to a rituximab-containing regimen. Patient characteristics are shown in Table 1. The median duration of study drug administration was 161 days (range, 5-1037 days), and the median number of cycles administered was 7 (range, 1-38).

3.2 | Safety and tolerability

Tirabrutinib was generally well tolerated (Table 2). The MTD was not reached. No DLT were observed in Cohorts 1-3. One DLT (grade 3 pneumonitis) was observed in 1 patient with WM in Cohort 4. After a 3-week drug-free period, the patient recovered with the administration of steroids and oxygen. The patient then continued treatment with tirabrutinib 160 mg QD and was undergoing treatment at the time of data cut-off (4 January 2018).

The most common AEs were rash (35.3%), vomiting (29.4%) and neutropenia (23.5%). Other AEs reported, for ≥15% of the patients, included: diarrhea (11.8%), skin disorder (23.5%), anemia (11.8%), headache (23.5%), and dyspnea (11.8%). No new safety concerns were identified.

### TABLE 3 Response rate by dosage of tirabrutinib

| Dosage            | N  | Overall response (CR, CRu, CRi, VGPR or PRa) | n (%) | [95% CI] |
|-------------------|----|---------------------------------------------|-------|---------|
| 160 mg QD         | 3  | 3 (100.0)                                   | [29.2, 100.0] |
| 320 mg QD         | 3  | 3 (100.0)                                   | [29.2, 100.0] |
| 480 mg QD         | 4  | 3 (75.0)                                    | [19.4, 99.4] |
| 300 mg BID        | 7  | 4 (57.1)                                    | [18.4, 90.1] |
| Total             | 17 | 13 (76.5)                                   | [50.1, 93.2] |

n (%) [95% CI] were estimated using the Clopper–Pearson method. CI, confidence interval; CR, complete response; CRi, CR with incomplete blood count recovery; CRu, CR unconfirmed; PR, partial response; VGPR, very good PR.

aIncludes patients with lymphocytosis assessed as modified PR.

**FIGURE 1** Maximum change in the orthogonal sum of the products of the greatest diameters of the target region (A) and the time to response and duration of response (B). +, censor; FAS, full analysis set.
population, included anemia, constipation, diarrhea, nausea, malaise, arthralgia and insomnia (17.6% each). Grade 3 or 4 AEs were mainly hematologic toxicities, with the most common being neutropenia (23.5%), anemia (11.8%) and leukopenia (11.8%). No AEs leading to death were reported.

Nine serious AEs (SAEs) were reported for 7 patients, of which 4 were reported as study drug-related, including Mallory-Weiss syndrome, de novo reactivation of HBV, acute myeloid leukemia and pneumonitis (1 patient each). One patient in the 160-mg QD cohort had de novo reactivation of HBV during Cycle 13; after starting entecavir, study drug treatment was continued. At screening, HBs antigen and HBV DNA tests were negative, and HBs and Hbc antibody tests were positive. Careful monitoring of liver function and HBV DNA was helpful to initiate the antivirus drug and continue the treatment of tirabrutinib. Mallory-Weiss syndrome was observed during Cycle 1 in 1 patient in the 480-mg cohort. Nausea and vomiting persisted in this patient after starting the study drug. No obvious hemorrhagic lesions were observed in the stomach or duodenum, but signs of Mallory-Weiss syndrome were observed in the esophagus. The patient received nothing by mouth and famotidine treatment was initiated. There was no tirabrutinib-induced lymphocytosis observed in this study.

3.3 | Efficacy

The ORR for all patients was 76.5% (13/17 patients) (Table 3), and the subgroup analysis showed that the ORR for non-GCB DLBCL was 75.0% (3/4 patients) (Table 4). Median progression-free survival in Cohort 3 was 109 days and was not estimable in Cohorts 1, 2 and 4. ORR was 80.0% (4/5 patients including 2 CR) for DLBCL, 40.0% (2/5 patients including 1 CR) for FL, 100% (4/4 patients including 3 CRs) for MCL, 100% (2/2 patients) for WM and 100% (1/1 patient) for CLL. The best overall response in DLBCL patients was CR in 2 patients and stable disease in 1 patient, among 3 patients who received 300 mg BID; in the 2 patients who received 480 mg QD, the best overall response was PR. Four patients with DLBCL (3/4 classified as non-GCB DLBCL) who achieved objective responses had no CD79A/B, MYD88 or CARD11 mutations. Of note, a CARD11 mutation was identified in 1 non-GCB DLBCL patient who did not achieve PR or a more response. At the time of the data cut-off, 1 CLL patient with a TP53 mutation (del 13q and del 17p) had achieved PR and was still receiving tirabrutinib therapy. Twelve of 16 patients with measurable lesions during the screening period showed ≥50% reductions in tumor diameter (Figure 1A). Of 13 patients who had achieved PR or more, 8 showed early response within 1 month after the beginning of the study treatment (Figure 1B).

CXCL-10 levels in responders tended to be lower on Day 8 of Cycle 1 (C1D8) than baseline and tended to be higher on Day 28 of Cycle 1 (C1D28) than on C1D8 (Figure 2). Among 13 responders, C1D28/C1D8 > 1 was reported in 8 (61.5%). Among 4 non-responders, C1D28/C1D8 > 1 was reported in 1 (50.0%) of the 2 with measurements available for C1D28. When patients with

TABLE 4 Subgroup analysis of overall response rate

| Diagnosis (subtype)         | n/N (%) [95% confidence interval] |
|-----------------------------|-----------------------------------|
| Non-GCB DLBCL               | 3/4 (75.0) [19.4, 99.4]           |
| DLBCL                       | 1/1 (100.0) [2.5, 100.0]          |
| Mantle cell lymphoma        | 4/4 (100.0) [39.8, 100.0]         |
| Follicular lymphoma         | 2/5 (40.0) [5.3, 85.3]            |
| Waldenström’s macroglobulinemia | 2/2 (100.0) [15.8, 100.0]     |
| Chronic lymphocytic leukemia | 1/1 (100.0) [2.5, 100.0]         |

The rates and their 95% confidence intervals were calculated using the Clopper-Pearson method.

DLBCL, diffuse large B-cell lymphoma; non-GCB DLBCL, non-germinat center B-cell-like diffuse large B-cell lymphoma.

CXCL-10 > 1000 pg/mL at baseline were excluded to eliminate the potential effect of inflammatory diseases, C1D28/C1D8 > 1 was reported in 7 patients out of 9 responders (77.8%). Among the 4 non-responders, 3 patients had baseline CXCL-10 levels > 1000 pg/mL, and in the remaining patient, CXCL-10 could not be measured on C1D28. There were no observable differences in the levels of other chemokines between responders and non-responders, including CCL-3, CCL-4, CXCL-4, CXCL-7, CXCL-12, CXCL-13 and BIM.
3.4 | Pharmacokinetics

The plasma concentrations of tirabrutinib for all cohorts are shown in Figure 3A,B. The PK parameters \( C_{\text{max}} \), \( T_{\text{max}} \), \( \text{AUC}_{12\text{h}} \), \( \text{AUC}_{24\text{h}} \), \( \text{AUC}_{\text{inf}} \) and \( T_{1/2} \) on C1D1 and C1D28 are shown in Table S1.

3.5 | Pharmacodynamics

Treatment with tirabrutinib suppressed the expression level of pBTK in all 11 patients with confirmed pBTK expression at the basal level. pBTK on C1D1 was rapidly eliminated in these 11 patients (data not shown). Expression was suppressed in the CLL patient in a time-dependent manner. Complete suppression was observed in the other patients at 2 hours post-dose on Day 1 of Cycle 1 (Figure 4). Basal expression of pBTK was below the detection limit in 6 of 17 patients.

4 | DISCUSSION

This study evaluated the MTD, safety, efficacy, PK and PD of tirabrutinib monotherapy in 17 Japanese patients with relapsed or refractory B-cell NHL and CLL. Furthermore, we conducted the first biomarker analysis of tirabrutinib in humans. We found that tirabrutinib was well tolerated and demonstrated promising efficacy in a group of Japanese B-cell NHL and CLL patients. Tolerability up to 300 mg BID was confirmed, and the MTD was not reached. Therefore, the recommended dosing schedule of tirabrutinib in Japanese patients with B-cell NHL and CLL is either 480 mg QD or 300 mg BID. However, it is still necessary to consider which dosing is recommended for each histological subtype.

No DLT was identified in Cohorts 1-3. A DLT (grade 3 pneumonitis) was observed in 1 WM patient in Cohort 4 (300 mg BID), but it resolved with supportive measures, and study treatment was resumed at a lower dose. Organized pneumonia was present before the patient entered the study. All AEs were manageable and resolved with temporary dose interruption or the administration of granulocyte colony-stimulating factor or other agents. There were no clinically significant AEs in this group of Japanese patients, and the safety profile and most common AEs (neutropenia and anemia) were similar to those reported in the European phase I study.

A response was seen beginning at 160 mg QD. No correlation was found between response rate and dose, possibly because the dose was gradually increased from an initial dose at which an adequate response was observed in a previous phase I study and the number of enrolled patients was small. Histological subtypes may have also influenced the response.
This study has some limitations, including the small sample size of each subtype. However, a notable result was that the ORR was 80% (4/5 patients) in patients with DLBCL. CR was seen in 2 patients in the 300 mg BID cohort, while the best response in the 480 mg QD cohort was PR. This trend toward improved response in the 300-mg BID cohort may be because the arithmetic mean trough tirabrutinib plasma concentrations on C1D28 were 3.2-fold higher with 300 mg BID than with 480 mg QD. This result is supported by pre-clinical pharmacological studies in the TMD8 model. Therefore, BID dosing may improve the efficacy of tirabrutinib in patients with DLBCL. Of note, 4 patients with DLBCL who achieved objective responses had no CD79A/B, MYD88 or CARD11 mutations. A study of DLBCL patients treated with ibrutinib reported that 55.5% of patients with ABC tumors with B-cell receptor mutations responded to treatment, and this rate increased to 80% for those who also had MYD88 mutations.

In the current study, 1 CLL patient (160 mg QD) had a TP53 mutation but achieved PR, and treatment is currently ongoing, despite the poor prognosis with TP53 mutation reported for this type of case treated by conventional chemotherapy. The efficacy of tirabrutinib against CLL with TP53 mutation may be comparable with that of ibrutinib.

For patients with CXCL10 levels > 1000 pg/mL, it was difficult to evaluate the effect of the study drug on CXCL10 levels because an increase in CXCL10 levels to >1000 pg/mL can be related to other complications (autoimmune disease or infection). Therefore, we only evaluated the change in CXCL10 levels in patients with CXCL10 ≤ 1000 pg/mL at baseline. CXCL10 levels in responders with CXCL10 ≤ 1000 pg/mL at baseline tended to be higher on C1D28 vs C1D8. This suggests that the administration of tirabrutinib suppresses CXCL10 production from tumor cells over a short period; therefore, a fast-acting antitumor effect rapidly lowers CXCL10 levels, and B-cell functions (including regulatory B-cells) are inhibited for a long period, accelerating Th1 functions.

Pharmacokinetics analysis demonstrated that the area under the concentration-time curve and maximum serum concentration were increased in a dose-dependent manner. Treatment with tirabrutinib at doses of ≥160 mg QD fully suppressed the expression of pBTK in all patients with confirmed basal expression of pBTK. In addition, pBTK in peripheral mononuclear cells was suppressed at 160 mg QD, indicating tumor shrinkage. However, although suppression of phosphorylation was seen in most patients, there were some non-responders. Therefore, further investigation is needed to validate pBTK as a biomarker.

In conclusion, in Japanese patients with B-cell NHL and CLL, tirabrutinib was well tolerated and showed promising efficacy with an acceptable PK profile for future development. Overall responses (≥PR) were observed in 76.5% of patients, including 4 DLBCL patients with no CD79A/B or MYD88 mutations, and 1 CLL patient with a TP53 mutation, providing promising data for future developments. Based on the results of this phase I study, phase II studies of tirabrutinib monotherapy have already been initiated in Japanese patients with specific subtypes of B-cell NHL. In addition, large-scale clinical trials of tirabrutinib are required to elucidate the differences between tirabrutinib and other BTK inhibitors, such as ibrutinib and acalabrutinib.

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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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