Association of immune-related pneumonitis with clinical benefit of anti-programmed cell death-1 monotherapy in advanced non-small cell lung cancer

Kana Ono | Hirotaka Ono | Yukihiro Toi | Jun Sugisaka | Mari Aso | Ryohei Saito | Sachiko Kawana | Tomoiki Aiba | Tetsuo Odaka | Suguru Matsuda | Shin Saito | Akane Narumi | Takahiro Ogasawara | Hisashi Shimizu | Yutaka Domeki | Keisuke Terayama | Yosuke Kawashima | Atsushi Nakamura | Shinsuke Yamanda | Yuichiro Kimura | Yoshihiro Honda | Shunichi Sugawara

Abstract

Background: The association between the development of checkpoint inhibitor pneumonitis (CIP) with tumor response and survival has remained unclear so far. The aim of the present study was to evaluate the association between CIP and the clinical efficacy of anti-programmed cell death-1 antibody in patients with advanced non-small cell lung cancer (NSCLC).

Methods: Between January 2016 and August 2019, 203 advanced NSCLC patients were administered with nivolumab or pembrolizumab. Comparisons were made between patients with and without CIP. We evaluated the time-to-treatment failure (TTF), progression-free survival (PFS), and overall survival (OS).

Results: CIP was observed in 28 (14%) patients. CIP was associated with a longer PFS (18.9 months [95% confidence interval, CI: 8.7 months–not reached] vs. 3.9 months [95% CI: 3.4–5.1 months, \(p < 0.01\)]) and longer OS (27.4 [95% CI: 20.7 months–not reached] vs. 14.8 months [95% CI: 11.2–17.9 months, \(p = 0.003\)]). Most patients discontinued the immune checkpoint inhibitor (ICI) treatment when they developed CIP. Seven patients (25%) lived for more than 300 days from treatment discontinuation and did not show any long-term tumor growth after treatment discontinuation.

Conclusion: CIP was associated with prolonged PFS and OS. Additionally, 25% of CIP patients did not show any tumor growth for long periods after treatment discontinuation. Careful management of CIP can help in obtaining the best clinical efficacy from anti-PD-1 antibody.

Keywords

anti-programmed cell death-1, checkpoint inhibitor pneumonitis, immune-related adverse events, non-small cell lung cancer, outcome
1 | INTRODUCTION

Lung cancer is the most common type of cancer-related death worldwide.1 Recently, programmed cell death 1 (PD-1) and programmed cell death ligand 1 (PD-L1) inhibitors (alone or in combination) have been shown to result in higher survival rates than standard chemotherapy in patients with advanced NSCLC.2-8 Administration of immune-checkpoint inhibitors (ICIs) is complicated by immune-related adverse events (irAEs) including pneumonitis, skin reactions, thyroid dysfunction, hepatitis, and infusion reaction; these differ from the adverse events of conventional systemic therapy.9 Some studies have reported that the occurrence of irAEs is linked to the clinical efficacy of ICIs in patients with NSCLC.10-12 However, it is unknown which irAE is particularly linked to the clinical benefit of ICIs.

Among individuals with irAEs, 5%–10% of the patients treated with ICIs developed checkpoint inhibitor pneumonitis (CIP), resulting in potentially serious toxicity.13 Similar to other irAEs, low-grade CIP was found in most cases, and it improved with immunosuppressive therapy. However, severe CIP can lead to fatal respiratory failure.14 The onset of CIP may be a reflection of the degree of immune activity, but it is currently unclear whether CIP development is an indicator of better antitumor response or clinical efficacy. We performed a retrospective study to investigate the association between CIP and the clinical efficacy of nivolumab or pembrolizumab in advanced NSCLC patients.

2 | MATERIALS AND METHODS

2.1 | Patients

We performed a single-institutional retrospective study of the medical records of patients with advanced NSCLC treated with either nivolumab or pembrolizumab between January 2016 and August 2019 at Sendai Kousei Hospital.

2.2 | Assessment

We analyzed CIP, skin reaction, infusion reaction, thyroid dysfunction, and hepatitis as irAEs. We defined irAEs as adverse events that require more frequent monitoring and may have an immunological basis requiring intervention with immunosuppression and/or endocrine replacement therapy.15 Patients were assigned to two groups (with or without CIP), and we evaluated their time-to-treatment failure (TTF), progression-free survival (PFS), and overall survival (OS). The best tumor response was defined with reference to the Response Evaluation Criteria in Solid Tumors (Ver. 1.1).16 We evaluated the clinical severity of irAE according to the Common Terminology Criteria for Adverse Events, version 4.0.

2.3 | Statistical analysis

Categorical variables were evaluated utilizing chi-square, Student’s t-test, Mann–Whitney U tests, or Welch’s t-test where appropriate. Survival outcome was estimated with the Kaplan–Meier curves and was compared between patient groups with the log-rank test. The relationship between patient variables and response was evaluated with univariate and multivariate logistic regression analyses. The Hazard ratios (HRs) were estimated using Cox proportional hazards. All p values were two-sided, and those <0.05 were considered statistically significant.

All statistical analyses were performed with EZR (Saitama Medical Center, Jichi Medical University, Saitama, Japan).17 This study was approved by the Sendai Kousei Hospital Institutional Review Board (IRB No. 30–36). Data were analyzed anonymously; therefore, there was no need to obtain informed consent from patients.

3 | RESULTS

3.1 | Patient characteristics

Two hundred and three patients (146 men [72%]; 57 women [28%]) received nivolumab (n = 141) or pembrolizumab monotherapy (n = 62) (Table 1). The median age was 70 (range 31–92) years. One hundred and ninety-five (96%) patients had an Eastern Cooperative Oncology Group Performance Status (PS) of 0 or 1. Eighty-one (40%) and 122 (60%) patients were diagnosed with squamous cell carcinoma and non-squamous NSCLC, respectively. Epidermal growth factor receptor

Lay summary

Checkpoint inhibitor pneumonitis (CIP) is one of the immune-related adverse events (irAEs). To date, an association between CIP and better tumor response or clinical outcome remains unclear. In our study, CIP was associated with progression-free survival (PFS) and overall survival (OS). Also, 25% of CIP patients did not have tumor progression long after treatment discontinuation. Most importantly, appropriate management was performed at the time of CIP onset. We think that careful management of CIP might maximize the clinical benefit of nivolumab and pembrolizumab monotherapy.
mutations were expressed in 21 (10%) patients. Thirty-eight (19%) patients had not been on any prior chemotherapy regimen, whereas 91 (45%), 37 (18%), and 37 (18%) patients had received 1, 2, or ≥3 courses of chemotherapy. PD-L1 was expressed in abundance (tumor proportion score ≥50%) in 51 (25%) patients, expressed at low levels (1% ≤ tumor proportion score <50%), absent (tumor proportion score <1%), and unknown in 152 (75%) patients. No patient had any active autoimmune disease and interstitial lung disease.

Complete response was seen in 3 (1%) patients, partial response in 57 (28%) patients, stable disease in 76 (38%) patients, and progressive disease in 67 (33%) patients. The objective response rate (ORR) was 30% (95% confidence interval [CI] 23–36), whereas the disease control rate was 67% (95% CI 60–73).

Immune-related adverse events.

The irAEs are summarized in Table 2. Of the 110 patients with irAEs, 58 (29%) patients presented with skin reactions, whereas 29 (14%), 28 (14%), 19 (9%), and 12 (6%) patients developed thyroid dysfunction, CIP, infusion reaction, and hepatitis, respectively.

The ORR was significantly better in patients who developed CIP and skin reactions than in those without CIP (68% vs. 24%, p < 0.001 and 53% vs. 20%, p < 0.001, respectively) (Table 2).

Table 3 shows the predictors of ORR, PFS, and OS in the multivariate analysis. CIP was also an independent predictor of ORR, PFS, and OS (ORR odds ratio 9.45 [95% CI: 3.35–26.6, p < 0.01]; PFS, HR 0.31 [95% CI: 0.18–0.56, p < 0.01]; OS, HR 0.31 [95% CI: 0.16–0.60, p < 0.01]).

Figure 1 shows a comparison of TTF, PFS, and OS for patients with and without CIP. CIP was associated with longer PFS (18.9 months [95% CI: 8.7–not reached] vs. 3.9 months [95% CI: 3.4–5.1 months, p < 0.01]). In addition, CIP was associated with longer OS (27.4 [95% CI: 20.7 months–not reached] vs. 14.8 months [95% CI: 11.2–17.9 months, p = 0.003]).

Figure 2 shows the PFS and OS by grade of CIP. The median PFS for patients with Grade 1 was 16.4 months (95% CI: 10–22.9 months) and 16.4 months (95% CI: 10–22.9 months).

| Variables          | n   | Onset time (week), Median | CTCAE ver. 4.0 Grade, n 1/2/≥3 | Tumor Response, n(%) CR/PR/SD/PD | Objective response rate (%) |
|--------------------|-----|--------------------------|--------------------------------|----------------------------------|----------------------------|
| Skin reaction      | 58(29%) | 6.4                      | 41/14/3                        | 2/29/23/4                        | 53%                        |
| Pneumonitis        | 28(14%) | 20                       | 5/16/7                         | 0/19/8/1                         | 68%                        |
| Infusion reaction  | 19(9%)  | 0                        | 13/6/0                         | 1/8/6/4                          | 47%                        |
| Thyroid dysfunction| 29(14%) | 7.8                      | 16/11/2                        | 0/13/10/6                        | 45%                        |
| Hepatotoxicity     | 12(6%)  | 8                        | 8/3/1                          | 0/5/6/1                          | 42%                        |

Abbreviations: CTCAE, Common Terminology Criteria for Adverse Events; CR, complete response; PR, partial response; SD, stable disease; PD, progression disease.
3.5–not reached), Grade 2 was 21.2 months (95% CI: 4.2–not reached), Grade 3 was not reached (95% CI: 2.0–not reached), and Grade 5 was 1.2 months. The median OS for patients with Grade 1 was 26.6 months (95% CI: 13.0–not reached), Grade 2 was 27.4 months (95% CI: 15.6–not reached), Grade 3 was not reached (95% CI: 6.3–not reached), and Grade 5 was 1.8 months.

The severity of CIP in response to ICI therapy. All patients with CIP are shown in Table 4. Of the 28 patients who had CIP, 7 developed Grade 3 or higher CIP, with good treatment responses (partial response or stable disease). All these patients with severe grade CIP were treated with steroid therapy for CIP, as indicated by the American Society of Clinical Oncology clinical practice guidelines. The cryptogenic organizing pneumonia (COP) pattern was the most frequent in 19 of 28 cases (68%), including the COP + ground-glass opacity (GGO) pattern.

Regarding the outcomes of CIP, there were 24 cases of improvement, 2 cases of no change, 1 case that worsened, and 1 case of death. Patient No. 25 died due to CIP; his chest

### Table 3

|                | ORR |          |          | PFS |          |          | OS  |          |
|----------------|-----|----------|----------|-----|----------|----------|-----|----------|
|                | Odds | 95% CI   | P        | HR  | 95% CI   | P        | HR  | 95% CI   |
| Sex (male)     | 0.86| 0.33–2.26| .77      | 1.08| 0.72–1.63| .70      | 1.28| 0.78–2.08| .33 |
| Age            | 1.00| 0.96–1.05| .97      | 1.00| 0.98–1.02| .90      | 1.01| 0.99–1.03| .29 |
| ECOG PS        | 0.75| 0.38–1.46| .40      | 1.18| 0.88–1.57| .26      | 1.81| 1.32–2.50| <.001|
| Smoking (current/past) | 2.2 | 0.59–8.25 | .24     | 0.87 | 0.54–1.41 | .57     | 1.10 | 0.62–1.95 | .75 |
| Prior chemotherapy regimens | 0.71 | 0.47–1.07 | .10     | 1.06 | 0.95–1.18 | .31     | 1.16 | 1.03–1.29 | .01 |
| Mutated EGFR, positive | 2.15 | 0.35–1.32 | .41     | 1.86 | 1.09–3.16 | .02     | 0.74 | 0.38–1.46 | .39 |
| Nivolumab/pembrolizumab | 0.99 | 0.41–2.43 | .99     | 1.02 | 0.70–1.48 | .93     | 1.19 | 0.77–1.84 | .44 |
| Skin reaction | 5.13 | 2.37–11.1 | <.001   | 0.40 | 0.27–0.58 | <.001   | 0.40 | 0.25–0.63 | <.001|
| Infusion reaction | 2.05 | 0.64–6.60 | .23     | 0.86 | 0.50–1.50 | .60     | 0.53 | 0.26–1.08 | .08 |
| Pneumonitis     | 9.45 | 3.35–26.6 | <.001   | 0.31 | 0.18–0.56 | <.001   | 0.31 | 0.16–0.60 | <.001|
| Thyroid dysfunction | 2.47 | 0.88–6.92 | .09     | 0.69 | 0.43–1.03 | .12     | 0.54 | 0.29–0.97 | .04 |
| Hepatitis       | 0.75 | 0.18–3.12 | .69     | 1.09 | 0.58–2.05 | .78     | 0.95 | 0.43–2.07 | .89 |

Abbreviations: irAE, immune-related adverse event; ORR, objective response rate; PFS, progression-free survival; OS, overall survival; PS, performance status; EGFR, epidermal growth factor receptor.

1 Results calculated with logistic regression.
2 Results calculated with Cox proportional hazard model.

**Figure 1** Time-to-treatment failure, progression-free survival, and overall survival in the study population. Kaplan–Meier curves are shown for time-to-treatment failure (A), progression-free survival (B), and overall survival (C) in patients with or without checkpoint inhibitor pneumonitis (CIP). The red line; with CIP; the black line; without CIP.
imaging showed worse findings, despite the administration of steroids. The mortality rate of all the CIP grades was 3.5%.

A swimmer plot displaying the course of treatment for all CIP patients is shown in Figure 3. Most patients discontinued ICI treatment when they developed CIP. Treatment of CIP was conducted in 24 of 28 (86%) patients: five patients received intravenous steroid pulse therapy, and 19 patients received prednisolone 0.5–2 mg/kg therapy. CIP was improved or resolved in 24 patients. Focusing on the period from ICI treatment discontinuation to disease progression or death, seven patients lived for more than 300 days, and these patients did not show any long-term tumor growth after treatment discontinuation (Figure 3 and Table 4; patient numbers 1, 2, 3, 4, 5, 6, and 7).

4 | DISCUSSION

4.1 | Overall result

In this study, we examined the correlation of CIP onset with tumor response and survival. A multivariate analysis revealed that CIP was associated with ORR, PFS, and OS. PFS and OS were longer in patients with CIP than in those without CIP, even though they had similar TTFs. To the best of our knowledge, this is the first report showing the development of CIP as an independent predictor of tumor response and survival in patients treated with anti-PD-1.

4.2 | IrAEs and tumor response

Some studies have reported that irAEs are related with better outcomes in melanomas.19-22 There are similar reports on lung cancer. We reported that the development of any irAE is associated with a longer PFS in patients with advanced NSCLC treated with anti-PD-1,10 and similar results were shown in other studies.11,12,23 For each irAE, we also reported that the development of skin reactions was correlated with a better response to anti-PD-1 antibodies.24 In addition, a past study reported that patients treated with pembrolizumab who developed immune-related thyroid dysfunction had a significantly longer OS than those who did not.25 These findings have led to reports linking the development of any irAE or a specific irAE with clinical benefits; however, it is unknown which irAE is particularly associated with clinical benefit. In this study, we found that CIP was a significant independent predictor of clinical benefit among the irAEs by multivariate regression.

4.3 | Checkpoint inhibitor pneumonitis

CIP is a significant adverse event that may lead to discontinuation of treatment and mortality. Symptoms of CIP are variable and nonspecific; the most common symptoms are dyspnea and cough, while fever and chest pain are less common. Furthermore, one third of patients may be asymptomatic at the onset.14 Therefore, members of the patient’s care team need to have a high level of awareness, so that changes in the early stages of the disease are not missed. Overall, the incidence of CIP is estimated to be between 3% and 6%.14 However, some reports showed a higher incidence between 13% and 19%, and our results are consistent with these reports.23,26-28 This difference in incidence may be due, in part, to the greater frequency of computed tomography (CT) in our routine clinical practice, which may have led to earlier detection, and to the inclusion of patients at potential risk for CIP who received PD-1 inhibitors outside of clinical trials.

Radiologic features of CIP were characterized into five subtypes: COP, GGO, interstitial lung disease, hypersensitivity, and pneumonitis not otherwise specified.14 Viral pneumonia such as coronavirus disease, alveolar hemorrhage, and interstitial pneumonia also show non-specific CT patterns.
### TABLE 4 All patients with checkpoint inhibitor pneumonitis

| Seq | TPS | ECOG PS<sup>a</sup> | Past regimens | Tumor response | CIP, Grade | Time to onset, days | Time to PD or cutoff from onset, days | Radiological pattern | Treatment | Outcome |
|-----|-----|----------------------|----------------|----------------|------------|----------------------|----------------------------------------|---------------------|-----------|---------|
| 1   | NA  | 1                    | 4              | PR             | 2          | 122                 | 1118                                   | COP                 | Prednisolone 1 mg/kg | Improved |
| 2   | 5   | 0                    | 1              | PR             | 3          | 141                 | 712                                    | COP                 | None       | Improved |
| 3   | 100 | 0                    | 1              | PR             | 2          | 202                 | 651                                    | COP                 | Prednisolone 0.5 mg/kg | Improved |
| 4   | 100 | 1                    | 0              | PR             | 3          | 149                 | 475                                    | NOS                 | None       | Improved |
| 5   | ≥75 | 0                    | 0              | PR             | 2          | 21                  | 341                                    | COP                 | Steroid pulse       | Improved |
| 6   | 100 | 0                    | 0              | PR             | 3          | 150                 | 329                                    | NOS                 | Prednisolone 1 mg/kg | Improved |
| 7   | NA  | 1                    | 2              | PR             | 2          | 561                 | 314                                    | COP+GGO             | Prednisolone 0.5 mg/kg | Improved |
| 8   | 10  | 1                    | 1              | SD             | 2          | 196                 | 279                                    | COP                 | Steroid pulse       | Improved |
| 9   | 0–24| 0                    | 1              | PR             | 1          | 42                  | 250                                    | COP                 | Prednisolone 1 mg/kg | Improved |
| 10  | ≥75 | 0                    | 0              | PR             | 2          | 21                  | 242                                    | COP                 | Prednisolone 2 mg/kg | Improved |
| 11  | 70–80| 0                   | 0              | PR             | 1          | 284                 | 214                                    | COP                 | Prednisolone 0.5 mg/kg | Improved |
| 12  | NA  | 0                    | 1              | SD             | 1          | 126                 | 212                                    | COP                 | None                  | Improved |
| 13  | ≥75 | 0                    | 1              | PR             | 2          | 175                 | 193                                    | COP                 | Prednisolone 1 mg/kg | Improved |
| 14  | 1–24| 0                    | 1              | PR             | 3          | 107                 | 159                                    | COP                 | Prednisolone 1 mg/kg | Improved |
| 15  | 0   | 0                    | 3              | SD             | 2          | 490                 | 155                                    | COP                 | Prednisolone 1 mg/kg | Improved |
| 16  | 50–74| 0                   | 0              | PR             | 3          | 141                 | 151                                    | GGO                 | Prednisolone 1 mg/kg | Improved |
| 17  | 0   | 1                    | 4              | PR             | 2          | 224                 | 142                                    | COP                 | Prednisolone 1 mg/kg | Improved |
| 18  | 60  | 0                    | 0              | SD             | 2          | 6                   | 123                                    | COP                 | Steroid pulse       | Improved |
| 19  | 1–24| 1                    | 2              | SD             | 2          | 35                  | 72                                     | interstitial        | Prednisolone 0.5 mg/kg | Improved |
| 20  | NA  | 0                    | 1              | PR             | 1          | 56                  | 51                                     | GGO                 | Prednisolone 1 mg/kg | Improved |
| 21  | ≥75 | 2                    | 0              | PD             | 2          | 16                  | 12                                     | GGO                 | Prednisolone 0.5 mg/kg | Improved |
| 22  | 100 | 0                    | 0              | PR             | 3          | 60                  | 3                                      | interstitial        | Prednisolone 0.5 mg/kg | Improved |
| 23  | 100 | 0                    | 0              | PR             | 2          | 857                 | 1                                      | COP                 | Prednisolone 0.5 mg/kg | Improved |
| 24  | 20  | 1                    | 1              | SD             | 2          | 166                 | 0                                      | COP                 | Steroid pulse       | Improved |
| 25  | 1–24| 0                    | 1              | SD             | 5          | 39                  | 0                                      | GGO                 | Prednisolone 1 mg/kg | Death       |
| 26  | NA  | 0                    | 3              | SD             | 1          | 791                 | 0                                      | COP                 | Steroid pulse       | No change   |
| 27  | NA  | 0                    | 1              | PR             | 2          | 581                 | 0                                      | COP+GGO             | None                | Improved |
| 28  | NA  | 1                    | 1              | PR             | 2          | 93                  | 0                                      | interstitial        | Prednisolone 0.5 mg/kg | No change |

Abbreviations: CIP, checkpoint inhibitor pneumonitis; COP, cryptogenic organizing pneumonia-like; ECOG PS, Eastern Cooperative Oncology Group performance status; GGO, Ground-glass opacities; NA, not assessment, NOS, Pneumonitis not otherwise specified; PD, progression disease; PR, partial response; SD, stable disease; TPS, Tumor proportion score.

<sup>a</sup>Scores range from 0 to 4, with high numbers indicating high disability.
We need to be careful in distinguishing these diseases from CIP. In the group with CIP, TTF was shortened, but PFS and OS were prolonged. However, in a previous report, OS was significantly shortened in patients with CIP. This difference may be in the PS variation. In our study, about 4% of all patients had PS 2–4, whereas they constituted 15.2% (26/170) of the sample size in a study by Fukihara et al. This difference in findings may be due to differences in patient characteristics.

Treatment of CIP is considered to discontinue the suspect drug for pneumonitis. Patients with drug-related pneumonitis also need to be treated with immunosuppressive drugs, but the response to systemic steroid varies with anticancer therapy. Some cases may be fatal despite receiving appropriate therapy (Table 4). In our study, the CIP mortality rate was 3.5% (n = 1). Previous studies have also revealed a CIP-related mortality rate ranging from 0% to 20%. The CIP-related mortality in patients treated with ICIs may be lower than that in patients treated with molecular targeted therapy or conventional chemotherapy. The different causes of pneumonitis, whether from ICI and other drugs, may have caused this difference. Pneumonitis after anticancer therapy may be caused directly by direct pulmonary toxicity or indirectly by activation of the body’s inflammatory response. Most conventional chemotherapy and molecular-targeted drugs directly injure airway epithelial cells, alveolar epithelial cells, and capillaries, resulting in non-reversible fibrosis. However, ICI treatment may lead to the over-activation of T cells that cause a reversible immune response in the lungs. These observations suggest that anti-PD-1 antibody-induced pneumonitis might be manageable with immunosuppressive therapy.

Figure 3 shows a swimmer plot of patients with CIP. Some patients with pneumonitis showed long-term recurrence after treatment discontinuation. A subset of patients who responded to anti-PD-1 in previous studies reported long-term clinical benefit even after discontinuation of therapy. In this study, although OS in patients with CIP was longer than in those without CIP, the factors are not clear. This good outcome may be related to early diagnosis and treatment, which ensures proper management at a less severe stage. We have an irAE management team (frontline immunotherapy team [FIT]) that enables us to perform a thorough physical
examination and to report and treat even mild symptoms as early as possible.

Presently, the mechanism by which the antitumor effect persists even after interruption of treatment with ICI has not been clarified. Osa et al. assessed the time of maximum duration of antibodies on T cells and the relationship between this duration and residual therapeutic effect or potential adverse events. They reported that the binding of nivolumab to memory T cells in the blood was detectable more than 20 weeks after the last dose, irrespective of the number of nivolumab doses or subsequent treatments. This mechanism suggests that long-term antitumor effects may be sustained. Therefore, appropriate management of CIP may have a good and long therapeutic effect.

4.4 Limitations

There are a few limitations to our study. The study had a small sample size, was a non-randomized, single-center cohort, retrospective study. Also, the expression level of PD-L1 was not measured because of the commercial unavailability of diagnostic kits at the beginning of the study in Japan. Hence, we were not able to sufficiently evaluate the therapeutic effect of PD-L1 expression. Recently, the combination of chemotherapy and immunotherapy has become the norm, reducing the chances of treating patients on anti-PD-L1 monotherapy. However, the results of this study may be useful in predicting the clinical efficacy of chemotherapy–immunotherapy combination therapy or immunotherapy combination therapy.

5 CONCLUSION

In our study, CIP was associated with prolonged PFS and OS. Moreover, in 25% of patients with CIP, tumors did not grow long after treatment was discontinued. Most importantly, management was carried out at the time of CIP onset and we believe that careful CIP management, especially early detection and treatment, will ease the attainment of the maximum clinical efficacy from anti-PD-1 antibody monotherapy.

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CONFLICT OF INTEREST

Dr. Sugawara reports lecture fees from Ono Pharmaceutical, Bristol-Myers Squibb, MSD, AstraZeneca, Chugai Pharma, Nippon Boehringer Ingelheim, Pfizer, Taiho Pharmaceutical, Eli Lilly and Company, Novartis, and Kyowa Hakko Kirin. Dr. Toi reports lecture fees from Ono Pharmaceutical, Bristol-Myers Squibb, MSD. Dr. Saito reports lecture fees from Bristol-Myers Squibb. Dr. Domeki reports lecture fees from Ono Pharmaceutical, Bristol-Myers Squibb. Dr. Nakamura reports lecture fees from MSD.

AUTHOR CONTRIBUTIONS

Drs Ono K, Toi, and Sugawara had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: Ono K, Toi, and Sugawara. Acquisition, analysis, or interpretation of data: Ono K, Ono H, Toi, Sugawara, Sugisaka, Kawashima, Aiba, Kawana, Saito R, Asa, Odaka, Matsuda, Saito S, Narumi, Tsurumi, Shimizu, Domeki, Terayama, Nakamura, Yamanda, Kimura, and Honda. Drafting of the manuscript: Ono K, Toi, and Sugawara. Critical revision of the manuscript for important intellectual content: Ono K, Ono H, Toi, Sugawara, Sugisaka, Kawashima, Aiba, Kawana, Saito R, Asa, Odaka, Matsuda, Saito S, Narumi, Tsurumi, Shimizu, Domeki, Terayama, Nakamura, Yamanda, Kimura, and Honda. Statistical analysis: Ono K, Toi, and Sugawara.

ETHICAL APPROVAL FROM AN INSTITUTIONAL REVIEW BOARD

The study was conducted with the approval of the Institutional Review Board of Sendai Kousei Hospital (IRB No. 30–36).

DATA AVAILABILITY STATEMENT

All data generated or analyzed during this study are included in this published article.

ORCID

Kana Ono https://orcid.org/0000-0002-1399-5778
Jun Sugisaka https://orcid.org/0000-0002-1000-0777
Sachiko Kawana https://orcid.org/0000-0001-7146-6949

REFERENCES

1. Siegel RL, Miller KD, Jemal A. Cancer statistics, 2016. CA Cancer J Clin. 2016;66(1):7-30.
2. Reck M, Rodríguez-Abreu D, Robinson AG, et al. Pembrolizumab versus chemotherapy for PD-L1-positive non-small-cell lung cancer. N Engl J Med. 2016;375(19):1823-1833.
3. Brahmer J, Rodríguez-Abreu D, Robinson A, et al. OA 17.06 updated analysis of KEYNOTE-024: Pembrolizumab vs platinum-based chemotherapy for advanced NSCLC with PD-L1 TPS ≥50%. J Thorac Oncol. 2017;12(11):S1793-S1794.
4. Gandhi L, Rodríguez-Abreu D, Gadgeel S, et al. KEYNOTE-189 Investigators. Pembrolizumab plus chemotherapy in metastatic non-small cell lung cancer. N Engl J Med. 2018;378(22):2078-2092.
5. Hellmann MD, Ciuleanu T-E, Pluzanski A, et al. Nivolumab plus ipilimumab in lung cancer with a high tumor mutational burden. N Engl J Med. 2018;378(22):2093-2104.
6. Borghaei H, Paz-Ares L, Horn L, et al. Nivolumab versus docetaxel in advanced nonsquamous non-small-cell lung cancer. N Engl J Med. 2015;373(17):1627-1639.

7. Brahmer J, Reckamp KL, Baas P, et al. Nivolumab versus docetaxel in advanced squamous-cell non-small-cell lung cancer. N Engl J Med. 2015;373(2):123-135.

8. Rittmeyer A, Barlesi F, Waterkamp D, et al. Atezolizumab versus docetaxel in patients with previously treated non-small-cell lung cancer (OAK): A phase 3, open-label, multicenter randomised controlled trial. Lancet. 2017;389(10066):255-265.

9. Weber JS, Kähler KC, Hauschild A, et al. Management of immune-related adverse events and kinetics of response with ipilimumab. J Clin Oncol. 2012;30(21):2691-2697.

10. Toi Y, Sugawara S, Kawashima Y, et al. Association of immune-related adverse events with clinical benefit in patients with advanced non-small-cell lung cancer treated with nivolumab. Oncologist. 2018;23(11):1358-1365.

11. Haratan K, Hayashi H, Chiba Y, et al. Association of immune-related adverse events with nivolumab efficacy in non–small-cell lung cancer. JAMA Oncol. 2018;4(3):374-378.

12. Teraoka S, Fujimoto D, Morimoto T, et al. Early immune-related adverse events and association with outcome in advanced non-small cell lung cancer patients treated with nivolumab: A prospective cohort study. J Thorac Oncol. 2017;12(12):1798-1805.

13. Georgia G, Vasilios T, Elias K, et al. Immune checkpoint inhibitor-related pneumonitis. Respiratation. 2020;99(11):932-942.

14. Naidoo J, Wang X, Woo KM, et al. Pneumonitis in patients treated with anti-programmed death-1/programmed death ligand 1 therapy. J Clin Oncol. 2017;35(7):709-717.

15. Weber JS, Hodi FS, Wolchok JD, et al. Safety profile of nivolumab monotherapy: A pooled analysis of patients with advanced melanoma. J Clin Oncol. 2017;35(7):785-792.

16. Eisenhauer EA, Therasse P, Bogaerts J, et al. New response evaluation criteria in solid tumours: revised RECIST guideline (version 1.1). J Clin Oncol. 2009;27(18):3488-3492.

17. Kanda Y. Investigation of the freely available easy-to-use software ‘EZR’ for medical statistics. Bone Marrow Transplant. 2013;48(3):452-458.

18. Brahmer JR, Lacchetti C, Schneider BJ, et al. Management of immune-related adverse events in patients treated with immune checkpoint inhibitor therapy: American Society of Clinical Oncology Clinical Practice Guideline. J Clin Oncol. 2018;36(17):1714-1768.

19. Sanlunorenzo M, Vujic I, Daud A, et al. Pembrolizumab cutaneous adverse events and their association with disease progression. JAMA Dermatol. 2015;151(11):1206-1212.

20. Hua C, Bousserma C, Mateus C, et al. Association of vitiligo with tumor response in patients with metastatic melanoma treated with pembrolizumab. JAMA Dermatol. 2016;152(1):45-51.

21. Freeman-Keller M, Kim Y, Cronin H, et al. Nivolumab in resected and unresectable melanoma: Characteristics of immune-related adverse events and association with outcomes. Clin Cancer Res. 2016;22(4):886-894.

22. Nakamura Y, Tanaka R, Asami Y, et al. Correlation between vitiligo occurrence and clinical benefit in advanced melanoma patients treated with nivolumab: a multi-institutional retrospective study. J Dermatol. 2017;44(2):117-122.

23. Sato K, Akamatsu H, Murakami E, et al. Correlation between immune-related adverse events and efficacy in non-small cell lung cancer treated with nivolumab. Lung Cancer. 2018;115:71-74.

24. Aso M, Toi Y, Sugisaka J, et al. Association between skin re- action and clinical benefit in patients treated with anti-programmed cell death 1 monotherapy for advanced non-small cell lung cancer. Oncologist. 2020;25(3):e536-e544.

25. Osorio JC, Ni A, Chaft JE, et al. Antibody-mediated thyroid dysfunction during T-cell checkpoint blockade in patients with non-small-cell lung cancer. Ann Oncol. 2017;28(3):583-589.

26. Suresh K, Voong KR, Shankar B, et al. Pneumonitis in non-small cell lung cancer patients receiving immune checkpoint immunotherapy: incidence and risk factors. J Thorac Oncol. 2018;13(12):1930-1939.

27. Peng TR, Tsai FP, Wu TW. Indirect comparison between pembrolizumab and nivolumab for the treatment of non-small cell lung cancer: a meta-analysis of randomized clinical trials. Int Immunopharmacol. 2017;49:85-94.

28. Passiglia F, Galvano A, Rizzo S, et al. Looking for the best immune-checkpoint inhibitor in pre-treated NSCLC patients: an indirect comparison between nivolumab, pembrolizumab and atezolizumab. Int J Cancer. 2018;142(6):1277-1284.

29. Shi H, Han X, Jiang N, et al. Radiological findings from 81 patients with COVID-19 pneumonia in Wuhan, China: a descriptive study. Lancet Infect Dis. 2020;20(4):425-434.

30. Fujikura J, Sakamoto K, Koyama J, et al. Prognostic impact and risk factors of immune-related pneumonitis in patients with non-small-cell lung cancer who received programmed death 1 inhibitors. Clin Lung Cancer. 2019;20(6):442-450.

31. Ando M, Okamoto I, Yamamoto N, et al. Predictive factors for interstitial lung disease, antitumor response, and survival in non-small-cell lung cancer patients treated with gefitinib. J Clin Oncol. 2006;24(16):2549-2556.

32. Kudoh S, Kato H, Nishiwaki Y, et al. Interstitial lung disease in Japanese patients with lung cancer: a cohort and nested case-control study. Am J Respir Crit Care Med. 2008;177(12):1348-1357.

33. Cho JY, Kim J, Lee JS, et al. Characteristics, incidence, and risk factors of immune checkpoint inhibitor-related pneumonitis in patients with non-small cell lung cancer. Lung Cancer. 2018;125:150-156.

34. Cooper JA Jr, White DA, Matthey RA, et al. Drug-induced pulmon-ary disease. Part 1: Cytotoxic drugs. Am Rev Respir Dis. 1986;133(2):321-340.

35. Michot JM, Bigenwald C, Champiat S, et al. Immune-related adverse events with immune checkpoint blockade: a comprehensive review. Eur J Cancer. 2016;54:139-148.

36. Gettinger S, Horn L, Jackman D, et al. Five-year follow-up of nivolumab in previously treated advanced non-small-cell lung cancer: results from the CA209-003 study. J Clin Oncol. 2018;36(17):1675-1684.

37. Osa A, Uenami T, Koyama S, et al. Correlation between vitiligo occurrence and clinical benefit in advanced melanoma patients. JCI Insight. 2018;3(19):e59125

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