A phase III study of adjuvant chemotherapy in patients with completely resected, node-negative non–small cell lung cancer (JCOG 0707)

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ABSTRACT

Objective: To evaluate efficacy of S-1 (tegafur/gimeracil/oteracil), an active novel fluoropyrimidine, as compared to UFT (tegafur/uracil) as a postoperative adjuvant therapy in patients with node-negative non–small cell lung cancer (NSCLC).

Methods: Eligible patients had undergone complete resection of p-stage I (T1 with tumor diameter >2 cm or T2-N0M0 by 5th edition Union for International Cancer Control TNM) NSCLC, and were randomized to receive oral UFT 250 mg/m²/day for 2 years (Arm A) or oral S-1 80 mg/m²/day for 2 weeks with a 1-week rest period, for 1 year (Arm B). The primary end point was relapse-free survival (RFS), with 80% power and a one-sided type I error of 0.05.

Results: From November 2008 to December 2013, 963 patients were enrolled (Arm A: 482, Arm B: 481). Toxicities (hematologic/nonhematologic) of grade 3 or more were observed in 15.9 (1.5/14.7)% in Arm A, and in 14.9 (3.6/12.1)% in Arm B, respectively. At data cut-off in December 2018, the hazard ratio for RFS was 1.06 (95% confidence interval, 0.82-1.36), showing no superiority of S-1 over UFT. The hazard ratio of overall survival (OS) was 1.10 (95% confidence interval, 0.81-1.50). The 5-year RFS/OS were 79.4%/88.8% in Arm A and 79.5%/89.7% in Arm B, respectively. The original NSCLC accounted for 58%/53%, respectively, of the Arm A/Arm B OS events. Secondary malignancies were observed in 85 (17.8%) and 84 (17.8%) individuals in Arm A and Arm B, respectively.

Conclusions: S-1 was not superior to UFT as postoperative adjuvant therapy in node-negative NSCLC. Future investigation should incorporate identification of high-risk populations for recurrence. (JTCVS Open 2020;4:90-102)

CENTRAL MESSAGE
S-1 (tegafur/gimeracil/oteracil) is an active novel fluoropyrimidine, but S-1 as adjuvant therapy failed to improve the outcome of patients with node-negative non–small cell lung cancer after surgery.

PERSPECTIVE
Outcome of patients with node-negative lung cancer is very good with Japanese standard of care, with 5-year overall survival of nearly 90%. In addition, there were significant competing risks, such as noncancer deaths and secondary malignancies. Therefore, future research should incorporate identification of, and be focused on, the patient groups at high risk for disease recurrence.

See Commentaries on pages 103 and 105.
Postoperative adjuvant therapy remains controversial for patients with node-negative (N0) non–small cell lung cancer (NSCLC) after surgical resection. Meta-analysis of platinum-based adjuvant therapy suggested its benefit in hilar node-positive and mediastinal node-positive patients with NSCLC. However, no significant benefit was observed in patients with pathologic stage (p-stage) IB (N0 and tumor >3 cm), and there was even a trend toward harm in patients with p-stage IA (N0 and tumor 3 cm or less). 1 Although suggestions are made for the benefit of platinum-based adjuvant chemotherapy in patients with N0 and a tumor size >4 cm, they are based on unplanned subset analyses of randomized trials alone. 2,3

In Japan, a series of randomized trials suggested a survival benefit of adjuvant therapy with the oral drug tegafur/uracil, or UFT, in patients with N0 NSCLC after complete resection. 4-7 UFT is a fluoropyrimidine drug with inhibition of dihydro-pyrimidine dehydrogenase, or DPD, which is a rate-limiting enzyme in the catabolism of fluorouracil and is associated with drug resistance. Although UFT is only marginally effective in advanced NSCLC, with response rate of only 6% to 7%, 8,9 its efficacy for postoperative adjuvant therapy in NO NSCLC was documented by a large randomized study by Kato and colleagues 10 and confirmed by a meta-analysis by Hamada and colleagues. 10 The meta-analysis showed that UFT was beneficial in patients with tumor size >2 cm. No significant interaction was observed between histology and UFT effect.

Tegafur/gimeracil/oteracil, or S-1, is a novel, more potent DPD-inhibitory fluoropyrimidine. 11,12 S-1 is active against a broad spectrum of tumors, including NSCLC, with a single-agent response rate exceeding 20%. 13-15 Its original administration schedule was 4 weeks on and 2 weeks off, but subsequent studies reported more favorable toxicity files and drug adherence with a modified schedule of 2 weeks on and 1 week off. 16 Postoperative adjuvant therapy with S-1 for 1 year improved survival outcomes of patients with gastric cancer (as compared with surgery alone) 17 or with pancreatic cancer (as compared to gemcitabine). 18 In addition, S-1 was superior to UFT as adjuvant therapy in terms of relapse-free survival in stage II/III rectal cancer. 19 Due to the mainly gastrointestinal toxicities of S-1, continuation of adjuvant therapy is hardly tolerable beyond 1 year, unlike UFT.

Based on the aforementioned rationales, we, Japan Clinical Oncology Group Lung Cancer Surgical Study Group (JCOG-LCSSG), conducted JCOG 0707 study, a randomized trial of adjuvant S-1 for 1 year versus standard UFT for 2 years, in patients with N0 NSCLC with tumor size >2 cm after complete surgical resection. The study objectives were to evaluate and safety of S-1 as compared with UFT to test the null hypothesis that there is no difference in relapse-free survival (RFS) between the arms. Since both belong to the same drug class, the clinical question should be, “does a more active (but toxic) drug do better” as adjuvant therapy for early-stage, N0 NSCLC. A part of the study results were presented at the World Conference on Lung Cancer on September 8, 2019 (Video 1).

METHODS
Study Design
This is an open-label, randomized phase III superiority trial in patients with N0 NSCLC after surgical complete resection to confirm the superior

VIDEO 1. The first author, Hideo Kunitoh, presented a part of the study results at the World Conference on Lung Cancer on September 8, 2019, at the mini-oral session “Challenges in the treatment of early stage NSCLC.” He emphasized the importance of “cost and value” issue at the conclusion of the presentation. The meeting was held in Barcelona, Spain, during September 7-10, 2019. Video available at: https://www.jtcvs.org/article/S2666-2736(20)30082-6/fulltext.
efficacy of adjuvant S-1 therapy over the standard of care, UFT adjuvant therapy. This study was compliant with the ethical principles of the Declaration of Helsinki. The trial protocol was approved by the institutional review boards of all participating institutions (approved by JCOG Committee Office on September 8, 2008). All patients provided informed consent in writing before enrollment. The trial was registered in the UMIN Clinical Trials Registry (UMIN000001494).

**Patients**

Patients were eligible for the study if they had histologically documented N0 NSCLC (based on the World Health Organization histologic classifications of 1999) and had undergone complete surgical resection within 56 days of enrollment. Computed tomography (CT) and/or magnetic resonance imaging of the brain or whole-body positron emission tomography (PET) scan was not mandatory at presurgical staging procedures.

Eligibility criteria also included (1) pathologic stage I, as defined by the 5th edition (1997) of Union for International Cancer Control TNM staging system; in cases with p-stage IA, the tumor diameter had to be more than 2 cm (T ≥ 2 cm). (2) Tumor resected with lobectomy, bi-lobectomy, or pneumonectomy, with lymph node dissection for N2a extent (ie, systemic hilar and mediastinal node dissection, excluding anterior and posterior nodes), or elective dissection. (3) No previous therapy except for surgical resection. (4) Eastern Cooperative Oncology Group Performance Status of 0 or 1 with adequate bone marrow, liver, and kidney function. (5) Oxygen saturation of 90% or more on ambient air. (6) Age 20 to 80 years. (7) Signed informed consent obtained from the patient.

The exclusion criteria and details of the eligibility criteria are provided in the Online Data Supplement.

On May 17, 2011, when 551 patients were randomized, patient enrollment was temporarily suspended following a report of the third treatment-related death in S-1 arm. The study was resumed on August 1, 2011, with a protocol amendment to tighten exclusion criteria. It was then specified that treatment-related death rate of 1% or more would be unacceptable in this good-risk patient group. The amended protocol also required strict informed consent, emphasizing the risks associated with S-1 therapy.

**Randomization and Masking**

Eligible patients were randomly assigned (1:1) to receive adjuvant S-1 therapy or UFT therapy. The JCOG Data Center performed this randomization using the minimization method to balance for institution, pathologic stage (IA or IB), histology (adenocarcinoma/not adenocarcinoma), age (below 65/65 or older), and sex (male or female). Investigators and patients were not masked to treatment allocation.

**Treatment**

In Arm A (UFT arm), oral UFT 250 mg/m²/day (tegafur dose) was administered in 2 or 3 doses per day, for 2 years. It was given continuously, without rest period. In Arm B (S-1 arm), Oral S-1 (Taiho Pharmaceutical Company, Tokyo, Japan) 80 mg/m²/day (tegafur dose) was administered in 2 doses per day continuously for days 1 to 14 followed by a rest period for days 15 to 21 (1 treatment cycle consists of 21 days), for 1 year.

Adverse events of the treatments were evaluated with the Common Terminology Criteria for Adverse Events, version 3.0. Treatment was suspended with the onset of clinically relevant toxicities and resumed upon recovery. The dosage was modified according to the criteria in the attached protocol.

In both arms, patients were regularly followed for the complete blood count, including neutrophil count, blood chemistry, oxygen saturation, physical examination, and drug adherence, as defined by the study calendar in the attached protocol. A chest radiograph was taken monthly for the first 3 months, tri-monthly thereafter up to 24 months, and every 6 months thereafter up until 60 months had elapsed. CT of the chest was obtained at 12, 36, and 60 months. Abdominal CT, brain CT and/or magnetic resonance imaging, bone scan, and/or PET scan were performed when disease recurrence was suspected. Details of treatments, their modifications, and clinical evaluation and monitoring schedule of the patients during and after the study therapy are provided in the Online Data Supplement. After completion/stoppage of the protocol treatment, patients should be followed with no additional therapy until disease relapse. Post-treatment after disease relapse is not regulated by the protocol, either after completion or stoppage of the protocol treatment. No crossover (S-1 therapy after UFT or vice versa) is allowed. Drug adherence was checked based on the self-report of each patient. Pill counts were not performed.

**End Points**

The primary end point was RFS for all randomized patients, which was calculated from the date of registration to the earliest date of disease recurrence or death from any cause. Differentiation of intrapulmonary metastasis (an RFS event) versus second primary lung cancer (not an RFS event) was made clinically by each attending physician, based on tumor histology, differentiation, and vascular invasion, etc. The primary end point was changed from overall survival (OS) to RFS, which was reported to be a surrogate for OS, by the protocol amendment in February 2014, as described in “protocol amendment” in the Appendix 1 and in the Online Data Supplement itself.

The major secondary efficacy end point was OS, which was measured from the date of registration to the date of death from any cause. Other secondary end points included pattern of recurrence, proportion of treatment completion, oral drug adherence, and the proportions of occurrence of adverse events, severe adverse events, and secondary malignancy.

**Statistical Analysis**

The planned sample size was 960 (expected total number of event of 302), which was determined using an accrual period of 5 years, a follow-up period of 5 years, a one-sided type I error of 0.05, power of 80%, 5-year RFS of 75% in UFT arm, and hazard ratio (HR) in the S-1 arm of 0.75 (5-year RFS of 80.6% in the S-1 arm).

A stratified log-rank test was performed in terms of RFS for all randomized patients using stratification factors selected from those used for randomization (pathologic stage [IA/IB] and histology [adenocarcinoma/not adenocarcinoma]). RFS and OS were estimated using the Kaplan-Meier method. HRs and their confidence intervals (CIs) were estimated by the Cox proportional hazards model. As a sensitivity analysis, we analyzed RFS and OS in all eligible patients. More details of statistical analysis are given in the Online Data Supplement.

**RESULTS**

**Patients**

From November 2008 to December 2013, a total of 963 patients were enrolled and underwent randomization, 482 to Arm A (UFT) and 481 to Arm B (S-1). More than 5000 patients were screened for, but did not participate in, the trial. The reasons for non-enrollment and patterns of care of these “non-participants” were collected in an observational study Comprehensive Support Project for Oncology Research (CSPOR) LC-03 and reported elsewhere. Four patients in UFT arm were ineligible due to advanced (T4) stage, concomitant malignancy, different histology (salivary type), or incomplete lymph node dissection. Five patients in S-1 arm were ineligible due to advanced stage (one T3 patient and two hilar node-positive patients), concomitant malignancy, or incomplete lymph node.
dissection. Ineligibilities were revealed with additional pathologic reports after patients were registered and randomized. They were all included in the intention-to-treat efficacy and safety analyses. Five patients in UFT arm and 9 patients in S-1 arm withdrew consent and did not receive the protocol treatment. They were all included in the efficacy analysis, but the safety analysis was performed for a total of 949 patients (477 in UFT arm and 472 in S-1 arm), all of whom received the treatment. Figure 1 shows the patient flow diagram.

Table 1 summarizes the patient characteristics. Information on the molecular change of the tumors, such as epidermal growth factor receptor–activating mutation or anaplastic lymphoma tyrosine kinase gene fusion, was not collected.

**Efficacy**

At data cut-off in December 2018, 5 years after the accrual of the last patient, 121 patients in UFT arm and 127 patients in S-1 arm, respectively, died or had disease relapse. Median RFS was not reached in either arm, and the 5-year RFS was 79.4% (95% CI, 75.5%-82.8%) in UFT arm, and 79.5% (95% CI, 75.6%-82.9%) in S-1 arm (Figure 2). The HR of S-1 arm as compared with UFT arm, estimated by a stratified Cox proportional hazards model, was 1.057 (95% CI, 0.824-1.356) with a stratified, one-sided log-rank test P value of .6684.

At data cut-off, 77 patients in UFT arm and 85 patients in S-1 arm had died. Median OS was not reached in either arm, and the 5-year OS was 88.8% (95% CI, 85.6%-91.3%) in UFT arm, and 89.7% (95% CI, 86.6%-92.1%) in the S-1 arm (Figure 3). The HR of the S-1 arm as compared to UFT arm was 1.102 (95% CI, 0.810-1.500), with one-sided P = .7318.

There was a significant number of deaths due to causes other than the original lung cancer. In the UFT arm, the original lung cancers accounted for 45 (58%) of the 77 OS events; 30 (39%) died due to other causes, and the cause...
Pre-specified subset analyses for sex, age, smoking, stage, tumor side, lymph node dissection area, pleural invasion, and histology revealed no remarkable results; S-1 was not superior to UFT for either RFS or OS in each analysis (Figures 4 and 5).

Patterns of disease recurrence are summarized as the first relapse sites shown in Table 2. Relapse occurred predominantly as distant metastases, without major imbalances between the arms.

Post-protocol therapy provided for 86 and 91 patients in the UFT and S-1 arms, respectively. Three in the UFT arm and 11 in the S-1 arm received post-protocol therapy before documentation of disease recurrence, as deviation from the protocol. In the UFT arm, treatment included chemotherapy in 49, radiotherapy in 21, and other modalities in 21 patients; in the S-1 arm, the respective numbers were 54, 31, and 13. Some patients received more than 1 treatment modality.

Figure 6 summarizes the results. RFS and OS were better than expected and not different between the arms, with significant competing risks.

Safety
In the UFT arm, Grade 3 or greater toxicities were observed in 15.9% of the patients; 1.5% experienced hematologic, and 14.7% non-hematologic toxicities, respectively. In the S-1 arm, Grade 3 or greater toxicities were observed in 14.9% of the patients; 3.6% experienced hematologic, and 12.1% non-hematologic toxicities. Details of the toxicities according to the treatment arms are summarized in Table 3. There were 4 patients who died within the protocol, likely due to cardiovascular causes: 1 in the UFT arm and 3 in the S-1 arm. The death of the patient in the UFT arm was due to aortic dissection and adjudicated not to be treatment related, whereas the deaths of patients in the S-1 arm were adjudicated to be treatment-related, probably due to fluorouracil-induced cardiac ischemia.25

Toxicities of UFT were unexpectedly much more severe and frequent as compared with previous studies. In the study of Kato and colleagues, 7 grade 3 or more hepatic toxicities occurred in less than 1%; in the current study, it was 8%. Since they were generally transient, closer follow could have detected asymptomatic abnormalities. However, other toxicities also increased, and exact causes of the discrepancy with previous studies remain unknown.

Dose reductions due to treatment toxicities were performed in 96 (20.1%) patients in the UFT arm, and 190 (40.3%) in the S-1 arm, mainly due to gastrointestinal adverse events. Adherence to treatment is summarized in Table 4; 287 patients (59.5%) in the UFT arm and 263 (54.7%) in the S-1 arm completed the protocol treatment.

During the follow-up period, secondary malignancy developed in 85 (17.8%) in the UFT arm and 84 (17.8%) in the S-1 arm. The most frequent primary site was lung

of death was unknown in 2 cases. In the S-1 arm, the original lung cancers accounted for 45 (53%) of the 85 OS events; 32 (38%) died due to other causes, 3 deaths were treatment related, and the cause of death was unknown in 5 cases.

**TABLE 1. Patient characteristics**

| Treatment arm | Arm A (UFT) (N = 482) | Arm B (S-1) (N = 481) |
|---------------|-----------------------|-----------------------|
| Sex           |                       |                       |
| Male          | 278                   | 279                   |
| Female        | 204                   | 202                   |
| Median age, y (range) | 65 (37-79)           | 66 (33-80)           |
| Median days from surgery to drug initiation (range) | 43 (14-66)          | 42.5 (16-67)         |
| Histology     |                       |                       |
| Squamous cell carcinoma | 69               | 75                    |
| Adenocarcinoma | 386                | 386                   |
| Large cell carcinoma | 9                | 9                     |
| Other         | 18                    | 11                    |
| ECOG PS       |                       |                       |
| 0             | 441                   | 428                   |
| 1             | 41                    | 53                    |
| Smoking habit |                       |                       |
| Never         | 193                   | 194                   |
| Ever          | 289                   | 287                   |
| Operation procedure |                 |                       |
| Pneumonectomy | 1                    | 1                     |
| Lobectomy     | 474                   | 472                   |
| Bi-lobeectomy | 7                    | 8                     |
| Comorbidities |                       |                       |
| Present       | 295                   | 277                   |
| Absent        | 187                   | 204                   |
| Tumor size, cm |                    |                       |
| <=3           | 260                   | 254                   |
| >3~<=4        | 140                   | 149                   |
| >4            | 82                    | 78                    |
| Pathologic T factor |                |                       |
| T1            | 218                   | 226                   |
| T2            | 263                   | 254                   |
| T3            | 0                     | 1*                    |
| T4            | 1*                    | 0                     |
| Pathologic N factor |            |                       |
| N0            | 482                   | 479                   |
| N1            | 0                     | 2*                    |
| Pathologic stage |                  |                       |
| Stage IA      | 218                   | 224                   |
| Stage IB      | 263                   | 254                   |
| Stage IIA     | 0                     | 2*                    |
| Stage IIB     | 0                     | 1*                    |
| Stage IIIB    | 1*                    | 0                     |

UFT, Tegafur/uracil; S-1, tegafur/gimeracil/oteracil; ECOG: Eastern Cooperative Oncology Group; PS, Performance Status; N0, node-negative; N1, hilar node-positive. *Ineligible.
in 81 patients, followed by stomach in 23, prostate in 12, colorectal in 12, breast in 9, and bladder in 8 patients.

**DISCUSSION**

Optimal management strategy for N0 NSCLC after surgical complete resection remains elusive. Although several guidelines approve the use of platinum-based adjuvant chemotherapy in large (>4 cm) tumors, they are based only on post-hoc subset analyses of randomized trials. While the recent revision of the staging classification of NSCLC upstaged large N0 tumors (>5 cm) to stage II or III, there is no direct evidence for the use of platinum-based chemotherapy for the N0 subset.

UFT is only marginally active against advanced NSCLC, but, when used in adjuvant settings, was reproducibly shown to improve the postoperative outcome of N0 NSCLC.
Although its effectiveness was not confirmed in the Western world, it is not that the effect was refuted by Western clinical trials; the trials were never conducted outside Japan. Without any other data against adjuvant UFT effectiveness, we have no other choice but to accept it as a control arm of our trial; a "surgery-only" arm was deemed ethically unacceptable.

S-1 is a newer drug in the same class, combining fluoropyrimidines and DPD inhibition. Its single-agent response rate against advanced NSCLC exceeds 20%\(^{13-15}\) versus less than 10% in UFT.\(^{5,9}\) In fact, postoperative adjuvant therapy with S-1 was reported to be superior to UFT with respect to RFS in rectal cancer.\(^{19}\)

However, we could not demonstrate the superiority of adjuvant S-1 therapy over UFT in N0 NSCLC. Although the number of events was less than planned even after protocol amendment, there was not even a trend suggesting S-1’s superiority. In addition, we could not find any subset, including larger tumors or specific histology, in which S-1 would be beneficial. S-1 was toxic, with 3 treatment-related deaths, although none occurred after the protocol amendment to tighten eligibility. The treatment strategy based on risk/benefit in adjuvant therapy for N0 disease could thus be quite different from that with a more advanced tumor; more toxic but active “new-generation drug” might not work well in this population.
Although our study yielded only negative results, and UFT is not implemented into practice outside Japan, our trial results could nonetheless give some insights and provide suggestions for future management of and research into N0 NSCLC.

The outcomes of patients in our trial were exceedingly good for both arms. Although our initial assumption was that the 5-year OS in the UFT arm would be 70% and that of S-1 would be 76.5% (HR, 0.75), these turned out to be nearly 90%. This finding is notable because we excluded patients with tumors of 2 cm or less. We must bear in mind that the prognosis of N0 NSCLC, even after excluding “very early” tumors, is excellent, and make future study plans accordingly.

In addition, there was significant competing risk in our study population, with about 40% of the deceased patients dying from causes other than the original lung cancer. There were thus even fewer disease-specific events. Future trials should therefore incorporate methods to predict who are likely to have disease recurrence. Liquid biopsy might be one method for the identification of these high-risk patients after apparently complete resection.29

Although our study yielded only negative results, and UFT is not implemented into practice outside Japan, our trial results could nonetheless give some insights and provide suggestions for future management of and research into N0 NSCLC.

![Figure 5](image-url)  
**Figure 5.** Subgroup analysis for overall survival. Forest plot for overall survival according to the potentially prognostic factors in the intention-to-treat population failed to detect any subgroup in which S-1 might be beneficial. S-1, Tegafur/gimeracil/oteracil; UFT, tegafur/uracil; CI, confidence interval; HR, hazard ratio; ND1, hilar lymph node dissection; ND2a, systemic hilar and mediastinal lymph node dissection excluding anterior and posterior mediastinal nodes; ND2b, systemic hilar and mediastinal lymph node dissection including anterior and posterior mediastinal nodes.
our “standard” UFT. With the refinement of the staging system that incorporate invasive tumor size, we should identify patients who could forgo any adjuvant therapy. Observational studies such as CSPOR LC-03 could be useful, and we are now analyzing the UFT effect in the real world. 30

The fact that 17.8% of the patients in each arm had secondary malignancies is of some concern, raising the possibility of carcinogenesis from fluoropyrimidine compounds. A previous study by Kato and colleagues7 reported that secondary cancers developed much less frequently, in 5% to 6% of the cases, and UFT did not appear to influence their development. In contrast, the CSPOR LC-03 observational study found that 797, or 13.5%, of the 5922 patients screened for JCOG0707 had concomitant malignancy and were excluded from the study.24 Therefore, it seems that secondary/concomitant malignancies are much more common in our recent patient population.

As in the report of Kato and colleagues,7 the most frequently observed secondary malignancy was lung cancer. It is often challenging to differentiate a secondary lung cancer from intrapulmonary metastasis. If more intrapulmonary metastases were misdiagnosed as secondary lung cancer than vice versa, the RFS would be overestimated. Future studies may have to incorporate molecular analysis for differentiation of secondary and recurrent tumors.

The strengths of our study are solid study design with randomization, large sample size, and the homogenous patient population of N0 NSCLC diagnosed by standard surgical procedures, with the exclusion of early tumors of 2 cm or less.

The limitations of our study include biologically unexplained (but clinically demonstrated) efficacy of the control UFT arm, better-than-expected patient outcomes leading to fewer events. In retrospect, we underestimated the OS of the control UFT arm, even if we excluded small, early tumors.

| TABLE 2. Initial relapse site |
|-----------------------------|
| Treatment arm               | Arm A (UFT) (N = 91) | Arm B (S-I) (N = 88) |
| Local                       | 6                  | 5                  |
| Hilar or mediastinal lymph node | 23                | 13                |
| Supraclavicular lymph node  | 6                  | 3                  |
| Pleura or pericardium       | 15                 | 14                 |
| Brain                       | 12                 | 18                 |
| Pulmonary metastasis        | 41                 | 37                 |
| Other distant metastasis    | 32                 | 19                 |

UFT, Tegafur/uracil; S-I, tegafur/gimeracil/oteracil. *A patient could have more than 1 site at the disease relapse, and the total number of patients exceeds 100% of the original population.

In this randomized phase III trial, adjuvant S-1 failed to improve the outcome of patients with node-negative, p-stage I NSCLC as compared with the standard UFT. The RFS/OS were better than expected in both arms, with significant competing risks. Future investigation should incorporate identification of the high-risk population.

Study design: Phase III randomized trial
Primary endpoint: RFS

1) No significant differences in RFS/OS.
2) RFS and OS: far better than expected in both arms.
3) Significant competing risks; further compromising statistical power.

Implications
1) Adjuvant UFT remains standard.
2) Future studies should incorporate identification of high-risk population.

FIGURE 6. In this randomized phase III trial, adjuvant S-1 failed to improve the outcome of patients with node-negative, p-stage I NSCLC as compared with the standard UFT. The RFS/OS were better than expected in both arms, with significant competing risks. Future investigation should incorporate identification of the population at high risk of recurrence. NSCLC, Non–small cell lung cancer; p-stage I, pathological stage I; UFT, tegafur/uracil; S-I, tegafur/gimeracil/oteracil; RFS, relapse-free survival; OS, overall survival, HR, hazard ratio; CI, confidence interval.
Competing risks such as deaths from other causes and secondary malignancies further compromised the statistical power.

Other weaknesses of our study design include the lack of masking, lack of uniform staging procedures including use of PET, and no pill counts. Information on detailed tumor size and adenocarcinoma subtype classifications is also lacking, since we used “old” classifications in the 1990s, which were widely used in Japan at the time of the study planning. Also missing are several prognostic factors, including pulmonary function data and detailed comorbidities.

In addition, information of the tumor molecular change, such as epidermal growth factor receptor mutation or anaplastic lymphoma tyrosine kinase infusion, was not collected in this trial. However, surgical resection remains the standard of care for early-stage NSCLC irrespective of its molecular marker, and the role of target-based adjuvant therapy is yet to be established, especially in N0 disease. Our results would give benchmark for future research, including trials with target-based drugs or immune-oncology agents.

In conclusion, postoperative adjuvant therapy with oral S-1 was not superior to that with UFT in patients with N0 NSCLC, and UFT remains the standard in this population. Future investigation should incorporate identification of the population at high risk of recurrence (Figure 6).

### TABLE 3. Toxicities of the therapy according to study arm

|                      | G1-2 (Arm A) (UFT) | G1-2 (Arm B) (S-1) | G3 (Arm A) (UFT) | G3 (Arm B) (S-1) | G4 (Arm A) (UFT) | G4 (Arm B) (S-1) | % Any (G1-4) | % G3/4 Missing |
|----------------------|-------------------|-------------------|-----------------|-----------------|-----------------|-----------------|--------------|--------------|
| Arm A                | 477               | 472               | 477             | 472             | 477             | 472             | 477          | 472          |
| Leukocytes           | 47                | 73                | 0               | 2               | 0               | 0               | 9.9          | 15.9         |
| Hemoglobin           | 197               | 341               | 1               | 0               | 0               | 1               | 41.5         | 72.5         |
| Platelets            | 240               | 230               | 1               | 0               | 0               | 0               | 50.3         | 48.9         |
| Neutrophils          | 175               | 209               | 6               | 14              | 0               | 1               | 37.9         | 47.5         |
| Non-hematologic      |                   |                   |                 |                 |                 |                 |              |              |
| Total bilirubin      | 143               | 166               | 0               | 0               | 0               | 0               | 30.0         | 35.2         |
| AST                  | 256               | 261               | 24              | 4               | 2               | 0               | 59.1         | 56.1         |
| ALT                  | 205               | 198               | 33              | 7               | 5               | 0               | 50.9         | 43.4         |
| Creatinine           | 101               | 76                | 0               | 0               | 0               | 0               | 21.2         | 16.1         |
| Fatigue              | 117               | 188               | 2               | 6               | 0               | 0               | 24.9         | 41.1         |
| Pigmentation         | 51                | 207               | *               | *               | *               | *               | 10.7         | 43.9         |
| Rash                 | 71                | 125               | 1               | 4               | 0               | 0               | 15.1         | 27.3         |
| Mucositis, oral      | 53                | 138               | 2               | 2               | 0               | 0               | 11.5         | 29.7         |
| Laryngeal mucositis  | 9                 | 21                | 0               | 0               | 0               | 0               | 1.9          | 4.4          |
| Anorexia             | 128               | 265               | 5               | 15              | 0               | 0               | 27.9         | 59.3         |
| Nausea               | 83                | 208               | 2               | 5               | 0               | 0               | 17.8         | 45.1         |
| Vomiting             | 29                | 68                | 0               | 0               | 0               | 0               | 6.1          | 14.4         |
| Diarrhea             | 56                | 145               | 4               | 0               | 0               | 0               | 11.7         | 31.6         |

G, Grade; UFT, tegafur/uracil; S-1, tegafur/gimeracil/oteracil; AST, aspartate transaminase; ALT, alanine transaminase. Asterisk indicates no statistical comparisons were performed between the arms.

### TABLE 4. Adherence to the treatment

| Treatment arm | Arm A (UFT) (N = 482) | Arm B (S-1) (N = 481) |
|---------------|------------------------|-----------------------|
| Treatment not started | 5                      | 9                      |
| Treatment administered | 477                    | 472                    |
| Median days of treatment duration (25%-75%) | 728 (281-730) | 350 (119-369.5) |
| UFT administered for 1 y or more % (95% confidence interval) | 337/477 |
| S-1 administered for 0.5 y or more % (95% confidence interval) | 70.7 (66.3-74.7) |
| One or more dose reduction performed | 96 (20.1%) | 190 (40.3%) |
| Protocol treatment completed | 289 (60.0%) | 263 (54.7%) |

UFT, Tegafur/uracil; S-1, tegafur/gimeracil/oteracil. *Completion of 2 years of UFT (Arm A) or 1 year of S-1 (Arm B), with or without dose reduction.

Other weaknesses of our study design include the lack of masking, lack of uniform staging procedures including use of PET, and no pill counts. Information on detailed tumor size and adenocarcinoma subtype classifications is also lacking, since we used “old” classifications in the 1990s, which were widely used in Japan at the time of the study planning. Also missing are several prognostic factors, including pulmonary function data and detailed comorbidities.

In addition, information of the tumor molecular change, such as epidermal growth factor receptor mutation or anaplastic lymphoma tyrosine kinase infusion, was not collected in this trial. However, surgical resection remains the standard of care for early-stage NSCLC irrespective of its molecular marker, and the role of target-based adjuvant therapy is yet to be established, especially in N0 disease. Our results would give benchmark for future research, including trials with target-based drugs or immune-oncology agents.

In conclusion, postoperative adjuvant therapy with oral S-1 was not superior to that with UFT in patients with N0 NSCLC, and UFT remains the standard in this population. Future investigation should incorporate identification of the population at high risk of recurrence (Figure 6).
Conflict of Interest Statement

Hideo Kunitoh reports personal fees from Taiho, personal fees from Coviden Japan, personal fees from MSD (Merck Sharp and Dohme), personal fees from EA Pharma, personal fees from Daiichi-Sankyo, personal fees from Johnson and Johnson, personal fees from Boehringer-Ingelheim, personal fees from Eisai, personal fees from AstraZeneca, personal fees from Chugai Pharmaceutical Co., Ltd, personal fees from Bristol-Myers Squibb Japan, personal fees from MSD (Merck Sharp and Dohme) Japan, personal fees from Chugai Pharmaceutical Co., Ltd, grants from MSD (Merck Sharp and Dohme) Japan, and grants from Boehringer-Ingelheim Japan, personal fees from Daiichi-Sankyo, personal fees from Chugai Pharmaceutical Co., Ltd, personal fees from Medtronic Japan, personal fees from Teijin Pharma, personal fees from Bristol-Myers Squibb Japan, personal fees from MSD (Merck Sharp and Dohme) Japan, personal fees from Chugai Pharmaceutical Co., Ltd, and personal fees from Johnson and Johnson K. K. Medical Company, outside the submitted work. Masashi Tsuboi reports personal fees from AstraZeneca KK, personal fees from Eli Lilly Japan, personal fees from Boehringer-Ingelheim Japan, personal fees from Daiichi-Sankyo, personal fees from Chugai Pharmaceutical Co., Ltd, personal fees from Johnson & Johnson Japan, personal fees from Medtronic Japan, personal fees from Teijin Pharma, personal fees from Bristol-Myers Squibb Japan, personal fees from MSD (Merck Sharp and Dohme) Japan, personal fees from Chugai Pharmaceutical Co., Ltd, and personal fees from Johnson and Johnson K. K. Medical Company, outside the submitted work. Kenji Suzuki reports grants and personal fees from Ethicon, grants and personal fees from Medtronic, grants and personal fees from Takeda, personal fees from Hisamitsu Pharmaceutical Co., Ltd, and personal fees from Astellas Pharma, outside the submitted work. Shun-ichi Watanabe reported no conflicts of interest. The editors and reviewers of this article have no conflicts of interest.

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Key Words: non–small cell lung cancer, node-negative, adjuvant chemotherapy, tegafur/uracil, tegafur/gimeracil/oteracil
APPENDIX 1

Protocol Amendment

The initial primary end point was overall survival (OS), with the planned sample size of 960 in total (expected total number of deaths: 302), which was determined with an accrual period of 3 years and a follow-up period of 5 years, using a one-sided type I error of 0.05, power of 80%, 5-year OS of 70% in UFT arm, and a hazard ratio (HR) in the S-1 arm of 0.75 (5-year OS of 76.5% in S-1 arm).

Monitoring in June 2013, after the completion of patient accrual, showed that the combined OS of the 2 arms was far better than expected, with 4-year OS of 91.6%. The original design was thus judged to be underpowered, and the study protocol was amended on February 28, 2014. The primary endpoint was changed to relapse-free survival (RFS), which was reported to be a surrogate for OS. However, the planned sample size was not changed from a total of 960 (expected total number of event of 302), which was re-calculated and determined using an accrual period of 5 years, a follow-up period of 5 years, a one-sided type I error of 0.05, power of 80%, 5-year RFS of 75% in UFT arm and HR in the S-1 arm of 0.75 (5-year RFS of 80.6% in the S-1 arm).