

Impact of select actionable genomic alterations on efficacy of neoadjuvant immunotherapy in resectable non-small cell lung cancer

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ABSTRACT

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Dr William N William Jr.; william.william@medicos oncoclinicas.com **Background** Neoadjuvant immune checkpoint inhibitors (ICIs) have improved survival outcomes compared with chemotherapy in resectable non-small cell lung cancer (NSCLC). However, the impact of actionable genomic alterations (AGAs) on the efficacy of neoadjuvant ICIs remains unclear. We report the influence of AGAs on treatment failure (TF) in patients with resectable NSCLC treated with neoadjuvant ICIs.

Methods Tumor molecular profiles were obtained from patients with stage I–IIIA resectable NSCLC (American Joint Committee on Cancer seventh edition) treated with either neoadjuvant nivolumab (N, n=23) or nivolumab+ipilimumab (NI, n=21) followed by surgery in a previously reported phase-2 randomized study (NCT03158129). TF was defined as any progression of primary lung cancer after neoadjuvant ICI therapy in patients without surgery, radiographic and/or biopsyproven primary lung cancer recurrence after surgery, or death from possibly treatment-related complications or from primary lung cancer since randomization. Tumors with AGAs (n=12) were compared with tumors without AGAs and non-profiled squamous cell carcinomas (non-AGAs+NP SCC, n=20).

Results With a median follow-up of 60.2 months, the overall TF rate was 34.1% (15/44). Tumor molecular profiling was retrospectively obtained in 47.7% (21/44) of patients and select AGAs were identified in 12 patients: 5 epidermal growth factor receptor *(EGFR)*, 2 *KRAS*, 1 *ERBB2*, and 1 *BRAF* mutations, 2 anaplastic lymphoma kinase *(ALK)* and 1 *RET* fusions. The median time to TF in patients with AGAs was 24.7 months (95% CI: 12.6 to 40.4), compared with not reached (95% CI: not evaluable (NE)–NE) in the non-AGAs+NP SCC group. The TF risk was higher in AGAs (HR: 5.51, 95% CI: 1.68 to 18.1), and lower in former/current smokers (HR: 0.24, 95% CI: 0.08 to 0.75). The odds of major pathological response were 4.71 (95% CI: 0.49 to 45.2) times higher in the non-AGAs+NP SCC

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Neoadjuvant immune checkpoint inhibitors (ICIs) with platinum-based chemotherapy are now a standard-of-care treatment for a subset of patients with operable non-small cell lung cancer (NSCLC). However, the effectiveness of ICIs in patients with tumors harboring select actionable genomic alterations (AGAs) remains unclear, particularly considering the poor response of epidermal growth factor receptor-altered and anaplastic lymphoma kinasealtered NSCLCs to ICI therapy in the metastatic setting.

WHAT THIS STUDY ADDS

⇒ We assessed the impact of select AGAs on the efficacy of neoadjuvant ICIs in patients with NSCLC treated with neoadjuvant nivolumab or nivolumab plus ipilimumab followed by surgery. Patients with tumors harboring AGAs exhibited higher rates of ICI treatment failure (TF), higher median percentage of residual viable tumor and shorter median time to TF.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ Our results suggest that certain AGAs may adversely impact the benefit of neoadjuvant ICIs in resectable NSCLC and underscore the importance of tumor molecular testing prior to neoadjuvant immunotherapy.

group, and the median percentage of residual viable tumor was 72.5% in AGAs compared with 33.0% in non-AGS+NPSCC tumors.

Conclusions Patients with NSCLC harboring select AGAs, including *EGFR* and *ALK* alterations, have a higher risk for TF, shorter median time to TF, and diminished pathological regression after neoadjuvant ICIs. The

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suboptimal efficacy of neoadjuvant chemotherapy-sparing, ICI-based regimens in this patient subset underscores the importance of tumor molecular testing prior to initiation of neoadjuvant ICI therapy in patients with resectable NSCLC.

INTRODUCTION

In operable non-small cell lung cancer (NSCLC), neoadjuvant immune checkpoint inhibitor (ICI) therapy is feasible and improves the rate of pathologic response when compared with standard-of-care chemotherapy.¹⁻⁴ Recently, the combination of chemoimmunotherapy received regulatory approval in the neoadjuvant and perioperative settings for a subset of patients with operable NSCLC based on the results of CheckMate-816, KEYNOTE-671, and AEGEAN trials.5-7 Major and complete pathologic response rates have been used as emerging candidate surrogate endpoints of clinical benefit after neoadjuvant therapies.⁸ However, little is known about the impact of genomic alterations on pathological responses and treatment failure (TF) after neoadjuvant ICIs.

The randomized NEOSTAR trial³ was a phase 2 study that reported on the outcomes of neoadjuvant ICI therapy using the programmed cell death protein-1 inhibitor nivolumab or the combination of nivolumab and the cytotoxic T-lymphocytes associated protein 4 inhibitor ipilimumab followed by surgery. Since the completion of the NEOSTAR trial, tumor molecular profiling is a highly recommended practice in the earlystage, operable setting for NSCLC. Adjuvant osimertinib, a treatment targeting epidermal growth factor receptor (EGFR) sensitizing mutations, is now the standard of care for patients with resected NSCLCs harboring these mutations.^{9 10} Likewise, in patients with anaplastic lymphoma kinase (ALK)-rearrangement positive NSCLC who have undergone surgical resection, adjuvant alectinib is now recommended.¹¹ Select phase 3 randomized controlled trials evaluating neoadjuvant and perioperative ICI therapy excluded patients whose tumors harbor EGFR and ALK alterations,^{5 7 12 13} and several ongoing trials are evaluating the role of perioperative targeted therapies for patients whose tumors harbor actionable genomic alterations (AGAs).^{14–16} This highlights the importance of tumor molecular profiling as an essential component of the decision-making process for neoadjuvant and perioperative immune-based treatment.

Patients treated on the randomized NEOSTAR trial received neoadjuvant nivolumab or nivolumab plus ipilimumab, irrespective of the presence of AGAs in their tumors, as testing of tumor molecular profiling was not mandated prior to initiation of neoadjuvant therapy in this study. With the availability of long-term follow-up data, we identified patients who experienced TF and explored the clinicopathological and molecular factors associated with TF. This exploratory analysis includes assessing the impact of AGAs on TF.

METHODS Patients and treatment

Patients with resectable stage I-IIIA (American Joint Committee on Cancer seventh edition staging system) NSCLC were randomized to treatment with either neoadjuvant nivolumab or nivolumab plus ipilimumab³ followed by surgical resection. The primary endpoint of the trial was the rate of major pathological response (MPR), defined as $\leq 10\%$ residual viable tumor in the (MPR), defined as ≤10% residual viable turnor in the original resected turnor after neoadjuvant therapy on the study.^{3 17} The clinical results have been previously reported.^{3 18} Patients underwent long-term follow-up for evaluation of survival outcomes.

was not required for trial enrollment. Post-hoc analysis of available tumor molecular profiling was performed for the current report. The specific profiling method depended on the availability and quality of appropriate tumor and/ or blood samples. Tumor samples, when available, underwent in-house next-generation sequencing (NGS assay, **f** DNA-based panel), capable of detecting 134 gene single nucleotide variants (SNVs) and 47 gene copy number gains, totaling 146 genes performed by the institutional Clinical Laboratory Improvement Amendments-certified went in-house next-generation sequencing (NGS assay, Molecular Diagnostic Laboratory at MD Anderson Cancer Center, as previously described.^{19 20} Additionally, targeted **6** intergenic and intragenic fusions in 51 genes were also text identified using an NGS assay (RNA-based panel). By clinician discretion and tissue availability, cytogenetic analyses by fluorescent in-situ hybridization (FISH) were performed to identify *MET* amplification and rearrangements in RET, ALK, and ROS1. In certain cases, in-house NGS-based circulating cell-free DNA analysis via institutional liquid biopsy was performed using a blood sample for genomic profiling, capable of detecting SNVs in 70 ≥ genes, 19 gene copy number gains, and six gene fusions, as ligand 1 (PD-L1) expression was evaluated on available 9 baseline tumor samples as part of the research correlative analyses, when possible, using single chromogenic immunohistochemistry analysis (clone 28-8, ab205921, 1:100 dilution, Abcam).³ The details of the tissue preparation process have been previously described.²² PD-L1-stained slides underwent scoring by two independent and trained pathologists following the guidelines recommended by the International Association for the Study of Lung Cancer.²³ Standard microscopy was used to identify any **8** positive membrane staining on malignant tumor cells and determine the tumor proportion score.²³ Tumor tissue molecular profiling reported here, involving NGS-based and/or cytogenetic (FISH-based) testing, was conducted on samples obtained from the primary tumor and/or surgical resection and/or metastatic lesion through bronchoscopic biopsy and/or radiographically guided percutaneous biopsy. Institutional liquid biopsy was obtained, where possible, through blood samples at the discretion

of treating physicians at the time of suspected or evident radiographic disease recurrence/progression during surveillance or on tissue confirmation of the presence of disease.

Current study objectives

The purpose of this exploratory report was to describe the impact of select AGAs on TF, MPR and tumor pathological regression in patients treated with neoadjuvant ICIs. Additionally, the study explored the impact of ICIbased therapy, targeted therapy, or chemotherapy, as a subsequent treatment option after failure from neoadjuvant ICIs. TF was defined as the progression of primary lung cancer after neoadjuvant therapy in patients without surgery, radiographic and/or biopsy-proven primary lung cancer recurrence after surgery, or death from possibly treatment-related complications or from primary lung cancer, from randomization on study. The database cutoff date was August 2, 2023.

Statistical methods

The association of MPR with molecular subgroups (AGAs vs non-AGAs+non-profiled squamous cell carcinomas (NPSCC)) was assessed by odds ratio in the univariable logistic regression model. The TF-free survival distributions were estimated by the Kaplan-Meier method. The univariable Cox proportional hazards (PH) regression model was used to evaluate the risk of TF for AGAs and non-AGAs+NPSCC groups, baseline clinicopathological characteristics and pathologic response in which the landmark method at surgery date was applied. The multivariable Cox PH regression model was also used to estimate the risk of TF for AGAs and non-AGAs+NPSCC groups. Comparison test was not performed as analyses were exploratory in nature. All statistics were performed in R (V.4.3.3), SAS (V.9.4), Excel (V.2016), and GraphPad Prism (V.10.0.3).

RESULTS

A total of 44 randomized patients who received at least one cycle of neoadjuvant ICI on trial were evaluated. With a median follow-up time of 60.2 months at the database cut-off, 15 patients (34.1%) had experienced TF. Among the 15 subjects with TF, there were a total of nine deaths, including 8 patients who had died after disease progression or recurrence, 1 patient died due to possibly treatment-related complications,³ 1 patient experienced surgery-precluding disease progression after neoadjuvant treatment and died of progressing disease after other therapies. Out of the 15 patients with TFs, there were 13 patients with documented recurrences, of which, 4 were locoregional and 9 were distant. Out of 15 TFs, 8 patients had received adjuvant chemotherapy and 6 had received postoperative radiotherapy. Among the 29 non-TF patients, 1 died from non-cancer cause, 1 from non-lung primary

cancer, and 2 from unknown causes without a known recurrence of their primary lung cancer (figure 1).

Characteristics of tumor AGAs and non-AGAs+NP SCC patients

Among the 44 patients, 21 (47.7%) had undergone tumor and/or blood molecular profiling. Of the 23 patients without profiling, 12 had adenocarcinomas and 11 had squamous histology at baseline (figure 2). In an exploratory fashion, we compared patients with AGAs (n=12) to the non-AGAs+NPSCC group (n=20), which consisted of two patient cohorts: Patients who underwent tumor molecular profiling but were found to lack AGAs (non-AGAs, n=9) and patients with NP SCC (n=11) (figure 2). ङ्

AGAs were found in 12 (57.1%) profiled patients. These consisted of five mutations in EGFR (2 Exon 21L858R mutations, 1 Exon 19 deletion with MET amplification, 1 Exon 19 deletion with RET deletion, 1 Exon 20 duplication with MET amplification), 2 KRAS G12C mutations, 1 ERBB2Y772 A775dup mutation, 1 BRAFV600E mutation, 2 ALK (1 EML4-ALK fusion, 1 rearrangement) fusions ٥ and 1 KIF5B-RET fusion. TP53 was the most frequently altered gene, present in 57.1% (12/21) of sequenced samples. The most frequent AGAs were EGFR mutations, which were found in 23.8% (5/21) of sequenced samples, followed by *KRAS* G12C mutations found in 9.5% (2/21). Of note, there were four *STK11* mutations, two of whom occurred with KRAS (1G12C, 1Q61H) co-mutation. Evalç uation of clinicopathological characteristics in AGAs and fe non-AGAs+NP SCC patient groups revealed that the AGAs group had more patients younger than 65, and more never-smokers. Non-AGA+NPSCC group had a higher data number of patients with squamous histology (table 1). Gender, race, clinical stage, and pretreatment tumor ining, PD-L1 expression status were not different between the two groups (table 1).

Clinicopathological and molecular characteristics of patients with TF

Al training, Among the 37 patients who were resected on-trial, 27.0% (10/37) had TF, and among the 7 patients who did not undergo surgical resection or were resected off-trial, <u>0</u> 71.4% (5/7) had TF. Out of 15 TFs, 60% (9/15) had 3 identifiable AGAs.

TF was more common in patients with AGAs than in the non-AGAs+NPSCC group (75% (9/12) vs 20% (4/20)). The most common AGAs in the TF group were EGFR **D** mutations (2 Exon 21 L858R mutations, 1 Exon 19 deletion with MET amplification, 1 Exon 19 deletion with RET **3** deletion, 1 Exon 20 duplication with MET amplification), followed by one EML4-ALK fusion, one KIF5B-RET fusion, one KRAS G12C mutation and one BRAF V600E mutation. Six squamous cell carcinomas were profiled, one had an ALK rearrangement, and five did not harbor an AGA.

Among the 32 patients in AGAs and non-AGAs+NP SCC groups, 28 had pathologic response assessment of residual viable tumor in the resected tumor specimen.



Figure 1 Treatment failure was defined as any progression of primary lung cancer in patients without surgery, radiographic and/or biopsy proven lung cancer recurrence after surgery, or death from possibly treatment-related complications or from primary lung cancer since randomization. Lung cancer recurrence/progression-free survival was defined as no evidence of progression and/or recurrence from the primary lung cancer at last radiographic scan since randomization. n, number.

The odds of achieving MPR in the non-AGAs+NPSCC group were 4.71 (95% CI: 0.49 to 45.2) times higher than in the AGAs group (figure 3A). The AGAs group

had a higher median percentage of residual viable tumor than the non-AGAs+NPSCC group (72.5% vs 33.0%) (figure 3A).



Figure 2 Patients with tumors harboring actionable genomic alterations (AGAs, n=12) were compared to the non-AGAs + NP SCC group, which included profiled tumors without AGAs and non-profiled squamous cell carcinomas (n=20); Tumor histology is at baseline. *Molecular profiling methods included next-generation sequencing (NGS DNA panel and/or NGS RNA fusion panel, when available), and/or cytogenetic studies (when available), and/or circulating cell-free DNA (ccfDNA) (when available) via liquid biopsy. ALK, anaplastic lymphoma kinase; EGFR, epidermal growth factor receptor; n, number; NP SCC, non-profiled squamous cell carcinomas.

Table 1	Clinicopathological characteristics of AGAs and
non-AGA	s+NPSCC patients

	AGAs (n=12)	Non-AGAs + NP SCC (n=20)	
Age			
< 65	7 (58%)	5 (25%)	
≥ 65	5 (42%)	15 (75%)	
Gender			
Female	4 (33%)	7 (35%)	
Male	8 (67%)	13 (65%)	
Smoking status			
Never smoker	5 (42%)	2 (10%)	
Former/current smoker	7 (58%)	18 (90%)	
Race			
White	9 (75%)	17 (85%)	
Non-white	3 (25%)	3 (15%)	
Stage			
Stage I/II	9 (75%)	15 (75%)	
Stage IIIA	3 (25%)	5 (25%)	
Histology*			
Squamous	1 (8%)	16 (80%)	
Non-squamous	11 (92%)	4 (20%)	
PD-L1†			
<1 %	6 (67%)	8 (67%)	
≥1 %	3 (33%)	4 (33%)	

Demographic information for patients with tumors harboring actionable genomic alterations (AGAs) or patients with tumors not harboring AGAs or non-profiled squamous cell carcinomas (non-AGAs+NP SCC). The percentages for each characteristic are rounded to the nearest whole number to total 100%. *Tumor histology at baseline.

[†]Pretreatment tumor PD-L1 expression status (n=21) on available samples by clone 28–8, Abcam^{3 23}; the PD-L1 expression status of three patients in AGAs and eight patients in non-AGAs+NP SCC were unavailable.

PD-L1, programmed death-ligand 1.

16.7% (2/12) of patients whose tumors had a *TP53* mutation had an MPR, and none (0/4) of the tumors harboring an *STK11* mutation achieved MPR. Three of the four patients with *STK11* mutations were in the non-AGAs+NPSCC group, and one (33.3%) had TF, this patient had a *KRAS* Q61H co-mutation. The only AGAs patients with MPR harbored a *BRAF* V600E mutation and developed recurrence in the brain 40.4 months after randomization and was treated with metastasectomy and postoperative radiation and remained disease-free at the time of dataset cut-off.

Patients with tumors harboring AGAs had a shorter median time to TF than those without AGAs. The median time to TF for AGAs patients was 24.7 months (95% CI: 12.6 to 40.4), and was not reached (95% CI: not evaluable

(NE)–NE) in the non-AGAs+NPSCC group (figure 3B). TF-free rates at 36, 48, and 60 months were 33.3% (95% CI: 10.3% to 58.8%), 22.2% (95% CI: 4.11% to 49.2%), and 22.2% (95% CI: 4.11% to 49.2%) for the AGAs group, and 78.8% (95% CI: 52.8% to 91.5%), 78.8% (95% CI: 52.8% to 91.5%), and 78.8% (95% CI: 52.8% to 91.5%) for the non-AGAs+NPSCC group.

The risk for TF was also assessed for association with molecular subgroup, age, sex, smoking status, race, clinical stage, tumor histology, MPR status (for resected patients ġ on-trial), and pretreatment tumor PD-L1 expression status (figure 4). The risk for TF was significantly higher for AGAs (HR: 5.51, 95% CI: 1.68 to 18.1), and lower in former/current smokers than never-smokers (HR: 0.24, 2 95% CI: 0.08 to 0.75). 80% (4/5) of never-smokers with **8** TF also had AGAs (2 EGFR, 1 EML4-ALK fusion, and 1 KIF5B-RET fusion). Patients with squamous histology also had a lower risk of TF compared with those with nonsquamous histology (HR: 0.20, 95% CI: 0.06 to 0.74). A multivariable Cox PH regression model was performed. However, since collinearity existed in histology and molecular subgroup, only smoking status and molecular subgroup were considered in the model. The AGAs had a higher risk for TF than the non-AGAs+NPSCC (HR: 4.35, 95% CI: 1.24 to 15.2) while the smoking status was no longer associated with TF (HR: 0.41, 95% CI: 0.12 to 1.37). The median time to first systemic retreatment was 20.2 months (range: 10.7-32.0) after randomization. The ð treatment course over time for non-AGAs+NP SCC (n=20, e patients 1-20) and AGAs (n=12, patients 21-32) patients is displayed in figure 5. The specific genomic alterations for the available patients are detailed in online supplemental table 1).

A sensitivity analysis was performed by assigning the two patients with KRASG12C to the non-AGAs+NP SCC group. The odds of achieving MPR in the non-AGAs+NPSCC + KRASG12C group were 3.38 (95% CI: 0.35 to 32.6) ≥ times higher than in the AGAs group (online supplemental figure 1A). The AGAs group had a higher median percentage of residual viable tumor than the non-g AGAs+NPSCC + KRASG12C group (61.0% vs 34.5%) (online supplemental figure 1A), AGAs had a median time to TF of 19.2 months (95% CI: 2.57 to 40.4), and was not reached (95% CI: NE-NE) in non-AGAs+NP SCC + KRASG12C group (online supplemental figure 1B). The technologies TF risk in the AGAs group was 5.96 (95% CI: 1.92 to 18.5) times higher than the non-AGAs+NPSCC + KRASG12C group.

Retreatment with ICI-based therapy

Among the 13 patients with TF and retreatment, 4 were retreated with ICI-based therapy, and 1 patient had disease control. This patient experienced disease recurrence 18.5 months after randomization and was treated with a chemoimmunotherapy regimen with stable disease for 2.1 months. Tumor profiling revealed *TP53* and *EGFR* L858R Exon 21 mutations while on ICI-based therapy and the treatment was subsequently changed to osimertinib





Figure 3 (A) Violin plot showing the pathological response in the resected tumor specimens from patients who underwent curative-intent surgery on trial in AGAs group (patients with tumors harboring an actionable genomic alterations, n=9) versus the non-AGAs+NP SCC group (patients with genomic profiling without AGAs or with non-profiled squamous cell carcinomas, n=19). The dds ratio of MPR was calculated in univariable logistic regression comparing the odds of achieving MPR in non-AGAs+NP SCC group with AGAs group. The dashed line indicates the median; the dotted lines indicate the lower quartile and upper quartile values; the top and bottom of the violin plots indicate the minima and maxima. (B) Kaplan-Meier curves of treatment failure-free survival probability for patients in the non-AGAs+NP SCC (n=20) and AGAs (n=12) groups. The 95% CIs are shown in parentheses by each treatment failure-free rate at 36, 48, and 60 months. The median time to TF with 95% CI was estimated using Kaplan-Meier method. Treatment failure was defined as any progression of primary lung cancer in patients without surgery, radiographic and/or biopsy-proven lung cancer recurrence after surgery, or death from possibly treatment-related complications or from primary lung cancer, since randomization. AGAs, actionable genomic alterations; *ALK*, anaplastic lymphoma kinase; *EGFR*, epidermal growth factor receptor; MPR; major pathologic response (residual viable tumor $\leq 10\%$ in the resected tumor specimen); NE, not evaluable; NP SCC, non-profiled squamous cell carcinomas; TF, treatment failure.

with disease control persisting 38.5 months after the first disease recurrence. The remaining three patients treated with ICI-based therapy did not achieve disease control. One patient recurred at 12.6 months and was treated with chemoimmunotherapy without response, while molecular profiling was pending. Molecular testing showed EGFR Exon 20 duplication with MET amplification and this patient was subsequently treated with targeted therapy with disease control for 6.20 months before radiographic disease progression. At the time of disease progression, treatment was changed to immunotherapy, but the patient died from lung cancer-related complications 2.47 months later. Another patient underwent definitive chemoradiation after neoadjuvant therapy (no surgery) and later experienced locoregional disease progression. Molecular profiling failed to reveal any AGAs and the patient started single-agent immunotherapy, but unfortunately died 4.80 months after treatment from cancer-related complications. Finally, one patient with recurrence 17.0 months after randomization was treated with one cycle of chemoimmunotherapy without response. The molecular profiling revealed a KIF5B-RET fusion, and the patient was enrolled in a RETTKI trial. However, the patient died

from disease-related complications 4.67 months after retreatment initiation.

Treatment with targeted therapy alone

In addition to the patients treated with ICI-based therapy and targeted therapy described above, four additional patients were treated with targeted therapy alone. One patient with EML4-ALK fusion had metastatic recurrence at 20.0 months after randomization and was treated with targeted therapy with disease control at 18.5 months until disease progression. One patient with EGFR Exon 19 deletion was treated with targeted therapy with disease control at 22.8 months of with ongoing treatment. One patient developed **g** recurrence at 15 months after randomization and B had EGFR L858R Exon 21 mutation, the patient was treated with targeted therapy with ongoing disease control at 27.5 months. Finally, one patient with EGFR Exon 19 deletion and MET amplification, had disease progression after neoadjuvant treatment, and was treated with definitive chemoradiation (no surgery). The disease eventually metastasized, and the patient was treated with targeted therapy, but ultimately died from complications secondary to lung cancer 6.37

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Figure 4 Forest plot showing the treatment failure HR for each variable. The HR from the univariable Cox proportional hazard regression estimates the risk of TF in the interest group relative to the risk in the reference group. Treatment failure was defined as any progression of primary lung cancer in patients without surgery, radiographic and/or biopsy-proven lung cancer recurrence after surgery, or death from possibly treatment-related complications or from primary lung cancer, since randomization. * Data collected at time of surgery in resected tumors in patients who underwent surgery on trial (n=28), three patients in AGAs and one patient in non-AGAs+NP SCC did not undergo surgery; time from surgery until treatment failure was used. ** Pretreatment tumor PD-L1 expression status (n=21) on available samples by clone 28-8, Abcam^{3 23}; the PD-L1 expression status of three patients in AGAs and eight patients in non-AGAs+NP SCC were unavailable. § Tumor histology at baseline. AGAs, patients with tumors harboring an actionable genomic alteration; MPR, major pathologic response (< 10% residual viable tumor in the resected tumor specimen); non-AGAs+NP SCC, patients with molecular profiling without AGAs or non-profiled squamous cell carcinomas; PD-L1, programmed death-ligand 1; TF, treatment failure; HR; hazard ratio,

months after targeted therapy initiation. In total, disease control was achieved with targeted therapy in 71.4% (5/7) of patients with TF who received targeted therapy alone or targeted therapy given in sequence with other systemic therapies.

KRAS and STK11 co-mutations

Among two patients with tumors harboring KRAS G12C mutations, one with STK11 co-mutations had locoregional recurrence 29.6 months after randomization and was treated with stereotactic body radiation therapy and the patient has had disease control for 14.4 months. The other patient with a tumor harboring KRAS G12C completed neoadjuvant ICI therapy followed by surgical resection on-trial and

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technol There were three additional patients who had tumors harboring an STK11 mutation. One patient with STK11 and KRAS Q61H mutations experienced disease recurrence 25.1 months after randomization. The patient was treated with chemotherapy regimens with disease control for 4.03 and 5.87 months but eventually experienced disease progression. The patient was then treated with a PARP inhibitor due to BRCA2 mutation for 2.03 months, but the disease progressed and chemoimmunotherapy plus bevacizumab was started with the completion of 2 cycles with stable disease at the time of data cut-off. Two other patients with STK11 mutation alone did not experience disease recurrence.



Figure 5 Swimmer plot showing duration of time free from treatment failure (blue) and time since treatment failure (red) in patients with tumors without an actionable genomic alteration (AGAs) or a non-profiled squamous cell carcinomas (non-AGAs+NP SCC, patients 1–20) and patients with tumor harboring an AGA (AGAs, patients 21–32). Chemotherapy, ICI-based therapy, and targeted therapy were started after treatment failure (TF). Non-profiled adenocarcinomas (n=12) are not included. Type of treatments each patient received are displayed chronologically based on time since randomization. Patient 23 had surgery off-trial. Patient 24 did not have disease progression after neoadjuvant therapy, was offered curative-intent surgery but declined. *ALK*, anaplastic lymphoma kinase; *EGFR*, epidermal growth factor receptor; ICI, immune checkpoint inhibitor.

DISCUSSION

In this study, we retrospectively evaluated the impact of select tumor molecular alterations on TF after neoadjuvant ICIs in patients with resectable NSCLC treated in a randomized phase 2 study. We found that patients with tumors harboring select AGAs had a greater median percentage of residual viable tumor after neoadjuvant ICI (chemotherapy-sparing) treatment when compared with those patients without AGAs in their disease. Patients in the AGAs group had a shorter median time to TF compared with those without these alterations. In aggregate, our results underscore the critical importance of tumor molecular profiling in patients with resectable NSCLC to guide the treatment decision-making process in order to maximize clinical outcomes.

Our findings suggest that neoadjuvant ICIs alone may not represent the most effective treatment approach for all patients. Most of the TFs were driven by patients with actionable tumor alterations including *EGFR*, *RET*, *KRAS* G12C, and *ALK*, among others, highlighting the need for pre-neoadjuvant therapy tumor molecular testing and appropriate targeted therapy-based approaches for select AGAs (including non-*EGFR* AGAs). Indeed, our study reveals that patients who experienced disease recurrence and were treated with appropriate targeted therapy achieved durable treatment response. In previous experiences with advanced and metastatic NSCLC, tumors harboring select driver alterations, including *EGFR* mutations and *ALK* fusions exhibited poor response to immunotherapy.^{24–26}

The appropriate use of therapies at the right treatment stage can provide durable benefits to patients. The field has been rapidly changing, with evidence that targeted therapy should be incorporated into the treatment paradigm in appropriate patients. For patients with *EGFR*-mutant resected NSCLC, the ADAURA trial⁹ ¹⁰

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demonstrated improved disease-free survival (DFS) and overall survival (OS) benefit with adjuvant osimertinib in stage IB-IIIA patients when compared with placebo. The ALINA trial¹¹ compared adjuvant alectinib to chemotherapy for stage IB-IIIA ALK+ NSCLC and demonstrated DFS benefit in both stage II-IIIA and intention-to-treat stage IB-IIIA groups, highlighting the benefit derived by tailoring targeted therapies to matched AGAs in the earlystage setting.

However, the effects of ICI therapy on patients with AGAs have been inconsistent in select trials in the earlystage setting. For example, the subgroup analysis of EGFR mutants from the AEGEAN trial evaluating perioperative durvalumab added to neoadjuvant chemotherapy versus chemotherapy plus placebo did not demonstrate eventfree survival (EFS) benefit at 12 or 24 months for EGFR mutant population, and with a 3.8% pCR rate compared with 17.2% in the overall population.⁷ On the other hand, subgroup analysis in the IMpower010 study²⁷ suggested a potential DFS benefit of adjuvant atezolizumab in EGFRpositive patients with PD-L1≥1%. EGFR and ALK-positive patients in KEYNOTE-671 also showed EFS benefit with perioperative pembrolizumab,⁶ however, it is worth noting that these observations were derived from a remarkably small sample size and should be interpreted with caution.

Furthermore, immunotherapy has been shown to improve OS in KRAS-mutants compared with chemotherapy,²⁸ however, STK11 co-mutation in KRAS-mutants has been associated with resistance to immunotherapy.²⁹³⁰ Due to the small sample size, our study does not allow for meaningful conclusions about the impact of KRAS co-mutations on the efficacy of neoadjuvant immunotherapy. Nonetheless, the present study extends the potential limited benefit from immunotherapy alone in patients harboring select (non-KRAS) AGAs in the neoadjuvant setting for early-stage resectable NSCLC. Integrating insights from other investigations, the optimal neoadjuvant treatment strategy for AGAs-associated NSCLC warrants further comprehensive exploration.

In the neoadjuvant setting, several efforts are ongoing to evaluate the role of targeted therapy with or without chemotherapy as compared with chemotherapy only. A phase 2 trial of neoadjuvant osimertinib (NCT03433469) demonstrated a 15% MPR rate and 48% partial response rate in appropriate stage I–IIIA operable patients.³¹ We await the results of the ongoing phase 3 NeoADAURA trial¹⁵ (NCT04351555), which is evaluating osimertinib with or without chemotherapy versus chemotherapy in patients with resectable EGFR-mutant NSCLC. The ongoing NAUTIKA-1 (NCT04302025) uses an umbrella design to screen for AGAs (ALK, ROS1, NTRK, BRAF, RET) and allocates patients to the appropriate neoadjuvant therapy trials prior to surgery.¹⁴¹⁶ This will provide further evidence for appropriate treatment strategy based on tumor molecular profiles. As the field advances, the timing and role of immunotherapy and targeted therapy with or without chemotherapy in the perioperative setting will need to be further defined.

Comprehensive tumor molecular testing can provide vital information for the treatment of NSCLC. Some guidelines suggest screening for select molecular markers (EGFR, ALK, ROS1, BRAF, MET, RET, ERBB2, KRAS) and evaluation of PD-L1 expression.³² Long turnaround time and lack of adequate tissue can pose logistical barriers, but despite these challenges, tumor molecular profiling should be performed, whenever feasible. This testing allows providers to tailor appropriate therapy to tumors harboring molecular alterations for which targeted **v** therapy has shown efficacy and is the standard of care, deal and to track progression of tumor biology for changes in driver alterations and development of therapeutic resistance.

Our study has several limitations. We recognize that 8 not all patients underwent tumor molecular profiling, which was due to the era of the trial initiation and rapidly evolving field and knowledge of resectable lung cancer. Notwithstanding, every effort was made to profile patient tumors when appropriate and feasible. Also, the NP SCCs were used in the comparison group since targetable genomic alterations are less common in squamous histology, though it is possible that these tumors also contained targetable alterations in the context of standard of care. Finally, it is important to note that this study was performed based on a single-center experience and is limited to a small sample size. Therefore, further validation in larger cohorts is warranted to strengthen the 5 text reliability and generalizability of our results.

In conclusion, for patients presenting with resectable and NSCLC and treated with neoadjuvant immunotherapy followed by surgery, tumors harboring AGAs portend a shorter median time to TF. This suggests that neoadjuvant immunotherapy alone may not be the most appropriate therapeutic option for select patients with targeted therapy-sensitive tumor alterations. The decision-making <u>></u> process leading to an evidence-based neoadjuvant treattraining, and similar technologies ment strategy should take several factors in consideration including the knowledge of molecular profiling of the otherwise operable NSCLC.

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Contributors NZ, WW, and TC designed the study. HL and JJL developed the statistical plan and supervised statistical analyses. NZ, CHL, HL performed all statistical analyses. NZ, CHL, AW, AP, MCBG, BWC, HTT, JW, LSS, HL, JJL, BS, and TC collected and/or analyzed the data. WW, FVF, AT, GRB, XL, JZ, FS, JMK, MA, CL, BSG, LB, YYE, RJM, DCR, GLW, WH, JAR, SGS, AAV, FS, DLG, JVH, BS, and TC recruited and/or treated patients. NZ, CHL, WW, HK, DLG, HL, JJL, JVH, MVN, BS, and TC interpreted the data. NZ, CHL, WW, HK, DLG, HL, MVN, BS, and TC wrote and/or edited the manuscript. All authors provided comments and approved the manuscript. TC is the guarantor.

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Supplementary Table 1		
Patient ID	MPR	Tumor moleculer profile
	status	i unior molecular prome
3 N	N- MDD	CG in baseline tumor:
	NO MPK	No alteration found
		NGS DNA in baseline tumor:
		PIK3R1 p.E291Q Exon 7 SNV Missense
		RAD50 p.S560C Exon 11 SNV Missense
c.		TP53 p.R175H Exon 5 SNV Missense
6	NO MPK	NGS RNA in baseline tumor:
		No fusion found
		ccfDNA for suspected recurrence:
		No alteration found
		NGS DNA in resected tumor:
		ARID1A p.V1585L Exon 18 SNV Missense
		FLT3 p.L860M Exon 21 SNV Missense
		NF1 p.P464L Exon 12 SNV Missense
		NF1 p.C2134F Exon 42 SNV Missense
8	No MPR	PMS2 p.A127T Exon 5 SNV Missense
		SMARCA4 p.R451L Exon 8 SNV Missense
		STK11 c.597+1G>T Splice Unknown
		TP53 p.Y236C Exon 7 SNV Missense
		NGS RNA in resected tumor:
		No fusion found
		NGS DNA in baseline tumor:
		FGFR3 Amplification chr4:1785640-1808399 4p16.3
		MYC Amplification chr8: 128748723-128753198 8q24.21
	No MPR	PTEN p.I135fs*44 Exon 5 Deletion Frameshift
10		TP53 p.V272M Exon 8 SNV Missense
		NGS RNA in baseline tumor:
		FGFR3-TACC3 fusion FGFR3-TACC3.F17T11.COSF1348
		CG in baseline tumor:
		No alteration found
		NGS DNA in resected tumor:
	No MPR	CDK12 p.A1098S Exon 12 SNV Missense
11		CDKN2A p.A100fs*46 Exon 2 Deletion Frameshift
		FBXW7 p.P230L Exon 4 SNV Missense
		NGS DNA in tumor at recurrence:
		CDK12 p.A10985 Exon 12 SNV Missense
		TP53 p.A78fs Exon 4 Duplication Frameshift
		TP53 p.L35fs Exon 4 Deletion Frameshift
		NGS RNA in resected tumor and tumor at recurrence:
		No fusion found

		NGS DNA in baseline tumor:
		ERBB2 p.T652M Exon 17 SNV Missense
14		NF1 p.S1443L Exon 32 SNV Missense
	MDD	NF1 p.S2460L Exon 50 SNV Missense
	MPK	NF1 p.K286E Exon 8 SNV Missense
		NOTCH3 p.V943L Exon 18 SNV Missense
		SETD2 p.K629M Exon 3 SNV Missense
		TP53 p.K291* Exon 8 SNV Nonsense
		NGS DNA in resected tumor:
		MYC Amplification chr8:128748723-128753198 8q24.21
		BRCA2 p.S2186fs*3 Exon 11 Insertion Frameshift
		FANCD2 p.I1249M Exon 37 SNV Missense
		KRAS p.Q61H Exon 3 SNV Missense
		MED12 p.K1225fs*74 Exon 26 Duplication Frameshift
		MYD88 p.I266fs*166 Exon 5 Insertion Frameshift
16		NF1 p.G2001V Exon 40 SNV Missense
16	NO MPK	PIK3R1 p.V589fs*13 Exon 14 Insertion Frameshift
		STK11 p.L282fs*5 Exon 6 Deletion Frameshift
		NGS RNA in resected tumor:
		No fusion found
		ccfDNA for suspected recurrence:
		No alteration found
		CG in resected tumor:
		No alteration found
		NGS DNA in resected tumor:
		NOTCH1 p.N2058fs*210 Exon 33 Duplication Frameshift
		NOTCH1 p.D1849H Exon 30 SNV Missense
10	No MPR	PTCH1 p.V1035L Exon 18 SNV Missense
10		STK11 p.Q37* Exon 1 SNV Nonsense
		TP53 c.919+1G>T Splice? Unknown
		CG in resected tumor:
		No alteration found
	MPR	NGS DNA in baseline tumor:
		CTNNB1 p.S37C Exon 3 SNV Missense
		NRAS p.Q61R Exon 3 SNV Missense
19		<i>RB1</i> p.Q344* Exon 10 SNV Nonsense
		TP53 p.V272L Exon 8 SNV Missense
		PIK3CA p.I391M Exon 7 SNV Missense
		CG in baseline tumor:
		No alteration found
		NGS DNA in tumor at progression:
21	No MPR	CDKN2A p.W15* Exon 1 SNV Nonsense
		EGFR Amplification chr7:55211009-55268091 7p11.2

		EGFR p.L747_T751del Exon 19 Deletion
		TP53 p.P222fs*25 Exon 6 Deletion Frameshift
		NGS RNA in tumor at progression:
		No fusion found
		ccfDNA at progression:
		CCND1 Amplification 11q13.3
		EGFR Amplification 7p11.2
		CDKN2A p.W15* Exon 1 SNV Nonsense
		EGFR p.L747_T751del Exon 19 Deletion
		TP53 p.P222fs*25 Exon 6 Deletion Frameshift
		CG in tumor at progression:
		<i>MET</i> amplification (<i>MET</i> copy number=10.60; MET:CC7 signal ratio = 2.20)
		NGS DNA in resected tumor:
		ATM p.V101L Exon 4 SNV Missense
		TP53 p.G244C Exon 7 SNV Missense
		NGS RNA in resected tumor:
22	No MPR	KIF5B-RET fusion KIF5B-RET.K15R12.COSF1232
		ccfDNA during confirmed recurrence:
		No alteration found
		CG in resected tumor:
		<i>RET</i> rearrangement
		NGS DNA in resected tumor:
		EGFR p.A767 V769dupASV Exon 20 Duplication
		NGS DNA in tumor at recurrence:
		EGFR p.A767_V769dupASV Exon 20 Duplication
		NGS RNA in resected tumor:
		No fusion found
		NGS RNA in tumor at recurrence:
		No fusion found
23	NO MPK	ccfDNA for suspected progression:
		NF1 Exon 18 Duplication Frameshift
		CG in baseline tumor:
		<i>MET</i> amplification (Copy number = 6.4; <i>MET</i> :CC7 signal ratio = 0.90)
		CG in resected tumor (ALK, ROS1 only):
		No alterations found
		CG in tumor at recurrence (<i>ALK, ROS1</i> only):
		No alterations found
		NGS DNA in baseline tumor:
		CCND3 Amplification chr6:41903565-41909230 6p21.1
	No MPR	NFE2L2 p.D29H Exon 2 SNV Missense
24		NOTCH1 p.M771I Exon 14 SNV Missense
		PIK3R1 p.H706P Exon 16 SNV Missense
		PMS2 p.G460C Exon 11 SNV Missense
	1	1 -

CG in baseline tumor: ALK rearrangement NGS DNA in baseline tumor: KRAS p.G12C Exon 2 SNV Missense TP53 p.S127F Exon 5 SNV Missense KIT p.M541L Exon 10 SNV Missense PIK3CA p.I391M Exon 7 SNV Missense CG in baseline tumor:	
ALK rearrangement NGS DNA in baseline tumor: KRAS p.G12C Exon 2 SNV Missense TP53 p.S127F Exon 5 SNV Missense KIT p.M541L Exon 10 SNV Missense PIK3CA p.I391M Exon 7 SNV Missense CG in baseline tumor:	
NGS DNA in baseline tumor: KRAS p.G12C Exon 2 SNV Missense TP53 p.S127F Exon 5 SNV Missense 25 No MPR KIT p.M541L Exon 10 SNV Missense PIK3CA p.I391M Exon 7 SNV Missense CG in baseline tumor:	
 <i>KRAS</i> p.G12C Exon 2 SNV Missense <i>TP53</i> p.S127F Exon 5 SNV Missense No MPR <i>KIT</i> p.M541L Exon 10 SNV Missense <i>PIK3CA</i> p.I391M Exon 7 SNV Missense CG in baseline tumor: 	
TP53 p.S127F Exon 5 SNV Missense25No MPR <i>KIT</i> p.M541L Exon 10 SNV Missense <i>PIK3CA</i> p.I391M Exon 7 SNV Missense CG in baseline tumor:	
25 No MPR <i>KIT</i> p.M541L Exon 10 SNV Missense <i>PIK3CA</i> p.I391M Exon 7 SNV Missense CG in baseline tumor:	
<i>PIK3CA</i> p.I391M Exon 7 SNV Missense CG in baseline tumor:	
CG in baseline tumor:	
No alteration found	
NGS DNA in resected tumor:	
CDK4 Amplification chr12:58142244-58145450 12q14.1	
MDM2 Amplification chr12:69203037-69233452 12q15	
26 No MPR NGS RNA in resected tumor:	
ALK EML4-ALK.E20A20 Fusion	
CG in resected tumor:	
ALK rearrangement	
NGS DNA in resected tumor:	
CDK4 Amplification chr12:58142244-58145450 12q14.1	
MDMR2 Amplification chr12:69203037-69233452 12q15	
<i>EGFR</i> p.L858R Exon 21 SNV Missense	
NGS RNA in resected tumor:	
No fusion found	
NGS DNA in resected tumor:	
CCND2 p.I163V Exon 3 SNV Missense	
KRAS p.G12C Exon 2 SNV Missense	
STK11 p.E57fs Exon 1 Deletion Frameshift	
NGS RNA in resected tumor:	
No fusion found	
CG in resected tumor:	
No alteration found	
NGS DNA in resected tumor:	
ERBB2 p.Y772_A775dupYVMA Exon 20 Duplication	
NGS RNA in resected tumor:	
No fusion found	
CG in resected tumor:	
No alteration found	
NGS DNA in resected tumor:	
MDM2 Amplification chr12:69203037-69233452 12q15	
30 No MPR EGFR p.E746_A750del Exon 19 Deletion	
POLE p.R231C Exon 7 SNV Missense	
NGS RNA in resected tumor:	
No fusion found	

		ccfDNA for suspected recurrence:
		No alteration found
		CG in resected tumor: 3' <i>RET</i> deletion
		NGS DNA in tumor at recurrence:
		EGFR p.L858R Exon 21 SNV Missense
2.1	No MPR	TP53 p.V274F Exon 8 SNV Missense
31		FGFR3 p.F384L Exon 9 SNV Missense
		NGS RNA in tumor at recurrence:
		No fusion found
32	MPR	NGS DNA in tumor at recurrence:
		ATR V1832I Exon 32 SNV Missense
		BRAF V600E SNV Missense
		CCND2 M196I Exon 4 SNV Missense
		MYCN E390K Exon 3 SNV Missense
		POLE Q352L Exon 11 SNV Missense
		RAD51D E50K Exon 3 SNV Missense
		NGS RNA in tumor at recurrence:
		No fusion found
Genomic altera	tions from I	NGS (next generation sequencing) platform and CG (cytogenetic) studies, and
circulating cell	free DNA (ccfDNA, next generation sequencing-based analysis performed on blood
sample via liqu	id biopsy).	MPR: major pathological response (viable tumor $\leq 10\%$ in resected specimen).
Splice?, an alter	ration of ge	nomic DNA adjacent to the intron/exon splice border which could have an
effect on RNA splicing has been identified.		