Neoadjuvant immunotherapy for non-small cell lung cancer: right drugs, right patient, right time?

Elizabeth Ahern, Ben J Solomon, Rina Hui, Nick Pavlakis, Ken O’Byrne, Brett M Hughes

ABSTRACT

Standard curative treatment of early-stage non-small cell lung cancer (NSCLC) involves surgery in combination with postoperative (adjuvant) platinum-based chemotherapy where indicated. Neoadjuvant immunotherapy (NAI) offers certain theoretical benefits compared with adjuvant approaches, including the ability to assess on-treatment response, reduce the tumor bulk prior to surgery, and enhance tolerability in the neoadjuvant setting. Indeed, the use of neoadjuvant therapies are well established in other cancers such as breast and rectal cancers to debulk the tumor and guide ongoing therapy, and neoadjuvant chemotherapy has similar efficacy but less toxicity in NSCLC. More recently, immune checkpoint inhibitors (ICIs) targeting programmed death-1 (PD-1)/PD-1-ligand 1 (PD-L1) have transformed the treatment of advanced NSCLC; the unique mechanisms of action of ICIs offer additional rationale for assessment in the neoadjuvant setting.

Preclinical studies in mouse cancer models support the proof of concept of neoadjuvant ICI (NAICI) through improvement of T-cell effector function and long-term memory induction. Preliminary early-phase human trial data support the proposition that NAICI in NSCLC may provide an feasible and potentially efficacious future treatment strategy and large, randomized phase III trials are currently recruiting to assess this approach. However, outstanding issues include defining optimal treatment combinations which balance high efficacy with acceptable toxicity, validating biomarkers to aid in patient selection, and avoiding potential pitfalls such as missing a window for successful surgery, that is, choosing the right drugs, for the right patient, at the right time. Predictive biomarkers to direct selection of therapy are required, and the validation of major pathological response (MPR) as a surrogate for survival will be important in the uptake of the neoadjuvant approach.

INTRODUCTION

Worldwide, lung cancer is the leading cause of cancer mortality with more than 1.7 million deaths in 2018 (18.4% total cancer deaths worldwide). Non-small cell lung cancer (NSCLC) accounts for most lung cancer diagnoses (84%). According to the Surveillance, Epidemiology and End Results database maintained by the National Cancer Institute (USA), 5-year survival for all patients with lung cancer is 19%. However, this varies by stage, with localized disease having 61% relative 5-year survival, falling to 35% for regional disease (spread to nearby structures or lymph nodes (LN)). NSCLC diagnosed at an advanced stage, where the cancer has spread to other parts of the body such as distant viscera, has a dismal 5-year prognosis at 6% survival.

Curative intent treatment approaches for early-stage disease often involve surgery, with other multimodality therapies recommended in certain settings to improve survival. Most commonly, chemotherapy and/or radiotherapy is administered in the adjuvant setting (postsurgery). Limited evidence suggests that chemotherapy given in the neoadjuvant setting (prior to surgery) is of comparable efficacy to adjuvant chemotherapy but with improved tolerability. Other proposed benefits of neoadjuvant chemotherapy (NACT) include increased likelihood to receive systemic treatment earlier to eradicate micrometastases, the ability to reduce tumor bulk prior to surgery potentially allowing more complete resections, and to directly observe the magnitude of pathological ‘in vivo’ regression which may aid in predicting subsequent outcome. On the other hand, a potential drawback of NACT is the risk of high-grade adverse events (AEs) which may impair the patient’s ability to proceed onto definitive surgery. This is an important consideration given that, for all the advances in systemic therapy, surgery remains the single most effective treatment modality contributing to cure in patients presenting with operable cancer.

More recently, immunotherapy comprising immune checkpoint inhibitors (ICIs) have changed the management of patients with advanced inoperable or metastatic NSCLC and has become part of the treatment paradigm for most patients with advanced disease...
Neoadjuvant approaches bring a unique opportunity to assess pathologic responses via definitive surgery. Despite OS remaining the gold-standard endpoint, pathologic response as a surrogate endpoint has been used in neoadjuvant trials in various cancer types. Pathological complete response (pCR), indicating no viable tumor remaining in the surgically resected specimen, has been correlated with OS in various malignancies following NACT, although the chance of achieving pCR is typically low. Adaptations of pathologic response rates include major pathological response (MPR), indicating ≤10% viable tumor cells remaining at surgical resection, or partial pathological response (pPR, ≤50% residual viable tumor), and these together with pCR have been variably assessed in NAICI trials in NSCLC and other cancers. MPR is prognostic in NSCLC following NACT as it is significantly correlated with improved survival. MPR has not yet been validated as a surrogate for survival in the setting of neoadjuvant immunotherapy, although MPR is accepted as a valid surrogate in neoadjuvant treatment of breast cancer, with Food and Drug Administration (FDA) approvals granted based on pathologic response rates for that disease.

PRECLINICAL RATIONALE FOR NAICI
Preclinical models of cancer allow for an examination of immunological mechanisms underpinning NAICI. NAICI has been explored in various orthotopic mouse models of cancer, where neoadjuvant compared with adjuvant immunotherapy approaches have resulted in improved outcomes. NAICI was also examined in a syngeneic subcutaneous model of NSCLC, where neoadjuvant combination anti-PD1 plus anti-cytotoxic T-lymphocyte-associated protein 4 (CTLA4) resulted in superior metastasis-free survival compared with the same combination immunotherapy given in the adjuvant setting. Preclinical studies further demonstrated some common T-cell-mediated mechanisms of anti-tumor activity. These included significantly enhanced influx of tumor-infiltrating lymphocytes (TILs), an improved tumor antigen-specific T-cell response within the primary tumor and the suggestion of enhanced long-term tumor-specific immunological memory. As will be discussed, these findings are reminiscent of those reported in human trials of NAICI in NSCLC.

A recurrent role of innate immune system elements such as innate inflammatory signaling pathways and intact type one interferon responses, together with the requirement for cross-presenting dendritic cells (DCs), has been noted. These suggest a potential benefit of T-cell priming, perhaps in the tumor bed or locoregional secondary lymphoid organs, as is being increasingly recognized as essential for effective immunotherapy. Analysis of contributions by innate immune elements and DCs have not yet been presented in human NSCLC NAICI studies, although in a neoadjuvant trial of immunotherapy in melanoma (NCT02437279), low baseline

MANAGEMENT OF EARLY-STAGE NSCLC
The improved survival for local and locoregional disease reflects that, in selected cases, management can be with curative intent—often with surgery as the mainstay of treatment. Surgery improves survival in early-stage disease: in a large retrospective cohort, those with stage I disease who were recommended but declined surgery had an estimated 5-year survival of 11%, compared with 54% in those who underwent surgery. However, fewer than 30% NSCLC patients receive surgery owing to factors as advanced tumor stage at presentation or medical comorbidities. Surgery remains the gold standard of treatment for medically and surgically appropriate stage I-II NSCLC patients: in stage I A-IB disease, 5-year overall survival (OS) post-lobectomy ranges from 45% to 65% and declines in a stage-dependent manner thereafter. Surgery is also recommended in selected patients with stage IIIA disease, often as part of multimodality therapy. Adjuvant chemotherapy is offered to patients with selected stage I-IIIA resected NSCLC and confers an additional absolute survival advantage of 5% at 5 years. The ADAURA trial reportedly demonstrated an impressive disease-free survival (DFS) benefit for osimertinib in resected epithelial growth factor receptor (EGFR)-mutated stage IB-IIIA NSCLC although OS outcomes are awaited.

NACT, given prior to surgery, may provide benefits such as improved tolerability, improved likelihood of receiving treatment and ability to reduce tumor size preoperatively. In trials comparing neoadjuvant against adjuvant or perioperative chemotherapy, rates of chemotherapy completion were higher in the neoadjuvant arms, although with no differences in toxicity. However, it is not clear if cumulative adjuvant chemotherapy dose is associated with improved survival in lung cancer. In two prospective NACT trials, a meta-analysis and a large retrospective study, the benefit of NACT in NSCLC appears similar or non-superior to that of adjuvant chemotherapy in terms of DFS. NACT can be part of the treatment paradigm for borderline-resectable or to downstage stage IIIA NSCLC patients as a bridge to surgery, for example in the presence of mediastinal LN metastases, although treatment decisions in this setting remain highly individualised.
tumorous expression of gene signature related to Batf3+ (cross-presenting) DCs correlated with risk of relapse.37

NEOADJUVANT IMMUNOTHERAPY TRIALS IN NSCLC

The ability to conduct 'window-of-opportunity' intervention studies pre-surgical resection is an attractive setting for translational studies into the mechanism and efficacy of NAICI. Studies involving NAICI in resectable NSCLC where some outcomes have been reported are summarized in table 1. The common findings include generally high resection rates, encouraging pathological regression rates and largely manageable toxicity profile.38–44 MPR has to date been the most common pathological response endpoint employed.

Forde et al published outcomes a in a phase II pilot study where 21 patients were treated with up to two doses of neoadjuvant nivolumab prior to planned resection of their tumors.38 Twenty-one patients were treated with nivolumab, but one was unresectable. The treatment was feasible and safe, without any treatment-related surgical delays, and with no previously unreported AEs. Only 23% patients had AE of any grade, with one event being grade 3 or higher (grade 3 pneumonia where surgery was subsequently successfully performed). At 1-year postsurgery, 80% of resected patients were alive and without tumor recurrence. Impressively, the MPR rate was 45% (9/20), including three patients with pCR in the tumor bed (although one of these still had small residual tumor in a hilar node). No patients had evidence of progression, with a median of 65% pathological regression in the primary tumor. Radiological response did not correlate with pathological response, with 90% patients appearing to have stable disease per Response Evaluation Criteria in Solid Tumors, perhaps reflective of the early timepoint of follow-up imaging necessitated by planned surgery for all patients 4 weeks after commencing nivolumab as well as tumor infiltration of immune cells. Predicted tumor-associated neoantigens (TANA) were determined through transcriptomic analysis. TANA-specific Tcells were shown in one patient to expand significantly by the time of the second dose of neoadjuvant nivolumab and enter a contraction phase by the time of surgery, recapitulating findings described above in preclinical models. Using multiplex immunohistochemistry (IHC), the authors reported influx of TILs (comprising lymphocytes and macrophages) in responding tumors.38 IHC has to date been the most common pathological response endpoint employed.

Table 1 Reported studies involving neoadjuvant immunotherapy in resectable non-small cell lung cancer

<table>
<thead>
<tr>
<th>Trial name/identifier</th>
<th>Trial design</th>
<th>Neoadjuvant trial intervention</th>
<th>Resection rate (%)</th>
<th>MPR rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOP1201, NCT01820754</td>
<td>Single arm, phase 2, n=24, Stage 1B-3A (18 with N2 disease)</td>
<td>Platinum-doublet NACT cycle followed by NACT plus ipilimumab (two further cycles)</td>
<td>54</td>
<td>NR</td>
</tr>
<tr>
<td>Forde et al, NCT02259621</td>
<td>Single arm, phase 2, n=21, Stage 1-3A</td>
<td>Nivolumab (two cycles)</td>
<td>95</td>
<td>45</td>
</tr>
<tr>
<td>NEOSTAR, NCT03158129</td>
<td>Randomized, phase 2, n=44 Stage 1-3A (single N2)</td>
<td>Arm A: nivolumab Arm B: nivolumab plus ipilimumab (three cycles)</td>
<td>83</td>
<td>Arm A: 22 Arm B: 38</td>
</tr>
<tr>
<td>LCMC3, NCT02927301</td>
<td>Single-arm, phase 2, n=181 Stage 1B-3B</td>
<td>Atezolizumab (two cycles)</td>
<td>88</td>
<td>18</td>
</tr>
<tr>
<td>Shu et al, NCT02716038</td>
<td>Single arm, phase 2, n=14 (of planned 30)</td>
<td>Platinum-doublet NACT+atezolizumab (four cycles)</td>
<td>87</td>
<td>57</td>
</tr>
<tr>
<td>NADIM Study-SLCG EudraCT: 2016-003732-20</td>
<td>Single-arm, phase 2, n=46 Stage 3A (N2)</td>
<td>Platinum-doublet NACT+neoadjuvant/adjuvant nivolumab (three neoadjuvant cycles)</td>
<td>89</td>
<td>83</td>
</tr>
<tr>
<td>NCT02904954</td>
<td>Randomized, phase 2, n=34 Stage 1-3A</td>
<td>Arm-1: durvalumab Arm-2: sub- ablative SBRT plus durvalumab</td>
<td>88</td>
<td>Arm-1: 0% Arm-2: 47%</td>
</tr>
</tbody>
</table>

*MPR rate based on all reported patients, not only on those who underwent resection. MRP, major pathological response; NACT, neoadjuvant chemotherapy; NR, not reported; SBRT, sub- ablative stereotactic radiotherapy.
cancer NAICI to date. Of the 181 reported patients, 159 had surgery (resection rate 88%) and MPR rate was reported as 21% (including 7% pCR), excluding those resected patients who were subsequently found to have oncogenic driver mutations or rearrangements (none of which had MPR). A low rate of grade 3–4 treatment-related AEs (TRAE) was reported preoperatively (6%) but two patients had grade 5 toxicity (both deemed non-treatment related). Interestingly, despite immature clonality and CD4+ effector memory T cell infiltration, CD8+ tissue resident and CD4+ effector memory T cells at surgery, and potentially greater induction of peripheral T-cell clonality and increased reponse with neoadjuvant NI as compared with N; these findings once again recapitulate those shown in preclinical models. At a median follow-up of 22 months, median recurrence-free survival and OS has not been reached in either cohort.

Chemotherapy and radiotherapy can enhance anti-tumor immunity in various ways, including by inducing immunogenic cell death and modulating immune cells in the tumor microenvironment. Neoadjuvant anti-PD1/PD-L1 chemoimmunotherapy in NSCLC has been explored in two phase 2 trials where data have been presented. MPR rates have been high (57%–74%). One trial enrolled 30 current or former smokers with stage IIB-III resectable NSCLC to receive up to four cycles of neoadjuvant atezolizumab plus carboplatin and nab-paclitaxel. Twenty-six participants successfully underwent R0 surgical resection, of which 14 had a thoracotomy. Of the 17 participants with MPR (57%), median DFS was 34.5 months compared with 14.3 months in those without MPR, but this did not reach statistical significance. Higher toxicity consistent with chemotherapy was seen, such as grade 3/4 neutropenia rates of 50%. Outcomes from the second trial, NADiM, assessing up to three cycles of neoadjuvant nivolumab plus carboplatin and paclitaxel followed by adjuvant nivolumab in 46 participants with resectable stage IIA NSCLC, are also impressive. A 12-month progression-free survival (PFS) (96%) and OS (98%) was reported. Forty-one (89%) participants had microscopically complete resection, and of those, at a median follow-up of 24.0 months, DFS was 85%. Postoperative surgical complications were noted in 12 of 41 (29%) resected participants such as infection or air leak. In this study, only 30% had grade 3 or higher toxicities of any type. Importantly, MPR achieved by 34 of 41 (83%) participants who underwent resection was significantly associated with improved 24-month PFS compared with those not achieving MPR in a post hoc analysis (88% vs 57%, p=0.01). Overall, 26 of 46 total trial participants achieved pCR. In both chemoimmunotherapy studies, no surgical delays were noted owing to neoadjuvant treatment. Finally, NCT02904954 explored two cycles of durvalumab with or without subablative radiotherapy. Overall, eight patients of 34 (24%) achieved MPR, all in the immunoradiotherapy arm. Preoperative high-grade toxicity rates (grade 3–4) were acceptable (12%). Addition of radiotherapy modulated the tumor microenvironment through influx of immune cells including lymphocytes and DCs.

Given these encouraging results, larger phase 3 randomized trials of neoadjuvant chemoimmunotherapy are currently recruiting (table 2). In these studies, the patient population has been refined to mainly include stage II–III (including some select IIIB) disease. In a press release, the trial sponsor has announced that CheckMate-816, assessing neoadjuvant nivolumab plus chemotherapy has met its primary endpoint of improved pCR compared with chemotherapy alone, but data are awaited.

**RIGHT DRUGS? EXPERIMENTAL STRATEGIES IN LUNG CANCER NAICI**

In addition to these ongoing neoadjuvant chemoimmunotherapy trials, ongoing smaller phase 1/2 studies of NAICI in resectable NSCLC are exploring novel combinations and targets, including combinational checkpoint immunotherapy, chemoimmunotherapy, immunoradiotherapy, and even chemoimmunoradiotherapy (online supplemental table 1). Overall, these studies have a variety of primary endpoints mainly relating to safety and feasibility and/or assessment of tumor response or survival outcomes. Furthermore, the patient populations are heterogenous, with selection of locoregionally advanced patients and/or potentially resectable patients among trials combining different modalities (eg, chemo-ICI or immunoradiotherapy), but with earlier stage patients permitted (stage I–IIIA) in the NAICI monotherapy trials. Two trials are assessing novel immunotherapy combinations in stage I–IIIA NSCLC patients (combining anti-PD1 with anti-Lag3 or anti-RANKL). Three studies are assessing neoadjuvant chemoimmunotherapy in stage III NSCLC patients, and these all include adjuvant ICI treatment postsurgery. Finally, two trials are assessing a potential role for neoadjuvant immunoradiotherapy in patients with stage III NSCLC, with one of these involving neoadjuvant chemoimmunoradiotherapy and adjuvant ICI.
Table 2  Ongoing phase 3 neoadjuvant chemoinmunotherapy trials in resectable non-small cell lung cancer

<table>
<thead>
<tr>
<th>Trial name/identifier</th>
<th>Trial design</th>
<th>Neoadjuvant trial intervention</th>
<th>Primary endpoint(s)</th>
<th>Estimated study completion (year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IMPower030, NCT03456063</td>
<td>Phase 3, two-arm, placebo-controlled, n=374 Stage 2, 3A, select 3B</td>
<td>Platinum doublet NACT±neoadjuvant/adjuvant atezolizumab</td>
<td>MPR, Event-free survival (EFS)</td>
<td>2024</td>
</tr>
<tr>
<td>CheckMate 816, NCT02998528</td>
<td>Phase 3, three-arm, open-label, n=642 Stage 1B-3A</td>
<td>Platinum doublet NACT±nivolumab vs nivolumab plus ipilimumab</td>
<td>EFS, pCR</td>
<td>2028</td>
</tr>
<tr>
<td>CA209-777, NCT04025879</td>
<td>Phase 3, two-arm, placebo-controlled, n=452 Stage 2-3B</td>
<td>Platinum doublet NACT±neoadjuvant/adjuvant nivolumab</td>
<td>EFS</td>
<td>2024</td>
</tr>
<tr>
<td>KEYNOTE-671, NCT03425643</td>
<td>Phase 3, two-arm, placebo-controlled, n=786 Stage 2, 3A, select 3B</td>
<td>Platinum doublet NACT±neoadjuvant/adjuvant pembrolizumab</td>
<td>EFS, OS</td>
<td>2026</td>
</tr>
<tr>
<td>AEGEAN, NCT03800134</td>
<td>Phase 3, two-arm, placebo-controlled, n=300 Stage 2, 3A, select 3B; EGFR/ALK wild-type</td>
<td>Platinum doublet NACT±neoadjuvant/adjuvant durvalumab</td>
<td>MPR</td>
<td>2024</td>
</tr>
</tbody>
</table>

ALK, anaplastic lymphoma kinase; EFS, event-free survival; EGFR, epithelial growth factor receptor; MRP, major pathological response; NACT, neoadjuvant chemotherapy; OS, overall survival; PCR, pathological complete response.

RIGHT PATIENT? BIOMARKERS OF EFFICACY IN NAICI

Limited translational data are available to inform appropriate patient selection for these various NAICI approaches in NSCLC, but are required to optimize efficacy while minimizing excess toxicity. Pretreatment biomarkers explored in advanced NSCLC, such as PD-L1 expression or tumor mutational burden (TMB), have shown heterogeneous associations with MPR rate in NAICI trials. PD-L1 expression was significantly associated with MPR rate in NEOSTAR but was not significantly associated with MPR in LCMC3, the Forde study, or the two chemoimmunotherapy studies including NADIM.38–40 In contrast, TMB was associated with MPR in Forde et al, but was not in LCMC3.38 39 More sophisticated potential biomarkers are suggested from some translational aspects of reported trials. In Forde et al, when comparing patients who achieved MPR versus no MPR, those with MPR had a more clonal T-cell population and a higher frequency of shared clones between tumor and peripheral blood.38 Overall tumor mutational burden and predicted TANA burden were both significantly correlated with degree of pathological tumor regression in this trial.38 Validating and integrating such investigations, which require sophisticated transcriptomic and in silico analysis, into the predictive algorithm when selecting appropriate patients for NAICI approaches at baseline is an emerging challenge.

Optimal methods of on-treatment efficacy assessment to guide clinical decision making also remain unclear. Changes in peripheral blood T-cell clonality after nivolumab treatment have been shown retrospectively to correlate with neoadjuvant nivolumab efficacy: increase in peripheral T-cells clones matching those within the tumor bed was correlated with pathological response.67 Assessment of response radiologically through cross-sectional imaging has not correlated pathologically with NAICI response.38 The PRADO trial in stage III melanoma hypothesizes that direct assessment of NAICI response in involved clinically evident LN via IHC may be used to de-escalate surgical therapy in patients achieving an MPR, building on previous findings in that disease of encouraging DFS post-NAICI for responders in that setting.35 68 However, preoperative sampling of tumor (primary or LN) has not been part of standard NSCLC response assessment after neoadjuvant approaches, and is a more complex procedure in thoracic malignancies; it has not been employed in NAICI NSCLC trials to date.

The approach to perioperative therapy in the setting of actionable molecular genomics findings (such as EGFR mutations or ALK-rearrangement) in resectable NSCLC requires more data to define optimal standard of care. In the adjuvant setting, long-term OS follow-up for third-generation EGFR tyrosine kinase inhibitor (TKI) osimertinib compared with chemotherapy is awaited from ADAURA, although a similarly designed study showing DFS benefit for the first-generation TKI gefitinib did not translate into an OS advantage.20 69 An adjuvant trial assessing alectinib in ALK-rearranged NSCLC is ongoing.70 A study assessing the neoadjuvant use of first-generation EGFR TKI erlotinib in potentially resectable stage IIIA NSCLC showed a significant improvement in PFS in the targeted therapy arm compared with chemotherapy, but a non-significant improvement in radiological response.71 The Lung Cancer Mutation Consortium LEADER trial aims to comprehensively assess neoadjuvant targeted therapies matched to oncogenic driver mutations in NSCLC (LCMC4). In contrast, PD1/PD-L1 pathway blockade has been associated with a lower chance
of immunotherapy benefit in metastatic EGFR-mutant or ALK-rearranged NSCLC, raising a concern about whether this is a suitable strategy in the neoadjuvant setting.40

RIGHT TIME? POTENTIAL PITFALLS OF IMMUNOTHERAPY GIVEN IN THE NEOADJUVANT SETTING

Optimal timing of immunotherapy in the treatment of early-stage NSCLC remains uncertain, although large adjuvant trials including ICI in the adjuvant treatment of NSCLC are ongoing with results due in the next few years (online supplemental table 2).73 Although theoretical benefits of NAICI relate to the mechanistic nuances of these therapies, certain pitfalls and hazards have been noted thus far in NAICI trials in cancer. First, a concern with the administration of systemic therapy in general is the risk of early progression of disease rendering the patient’s tumor inoperable. In LCMC3, 10 of 101 patients evaluable in the safety analysis were either found to be inoperable at exploration or had progression of disease during neoadjuvant atezolizumab resulting in abandonment of surgical plans.39 This trial permitted the enrolment of patients with stage IIIB NSCLC, and the proportion of those subsequently deemed inoperable being of more advanced stage is yet to be revealed. However, the prospect of progression during neoadjuvant treatment compromising surgical plans remains a concern. In a prior study, radiological reassessment immediately prior to surgery did not correlate with pathological response, so the optimal method of monitoring progress on NAICI is uncertain.38 Furthermore, in NEOSTAR, an ‘immune flare’ phenomenon was noted radiologically in LN in up to 11% patients where granuloma formation without tumor was found in the nodes pathologically. There might be a risk of avoiding curative surgery owing to concerns about disease progression.40

Second, although undue toxicity has not been noted in the NAICI NSCLC trials reported to date, the possibility that the host systemic immune system is more functional and less systemically suppressed in the setting of early compared with late cancer,36 74 potentially conveys the risk of marked immune-related-AE (irAE) development occurring concurrently with enhanced immune-mediated tumor regression. In an early NAICI trial of combination ipilimumab plus nivolumab in melanoma, 18 of 20 participants (90%) had grade 3/4 irAE noted, most of which (17 of 18) required cessation of therapy.96 Interestingly, this was reminiscent of a contemporaneous trial of preoperative combination ICI in resectable stage III or oligometastatic resectable stage IV melanoma where more than 70% patients had grade 3 or higher treatment-related TRAEs.25 The high irAE rate in early-stage melanoma in the setting of combination NAICI was subsequently shown to be likely dose-related, and adjustment of ipilimumab dose in particular is likely to have decreased irAE incidence while preserving efficacy in a follow-up trial.76

Surgical comorbidity arising from NAICI treatment is another potential concern. In the pilot study of neoadjuvant nivolumab in resectable NSCLC, 14/20 surgeries were performed with an open approach, including 7/13 which were converted from an initial minimally invasive approach, related in most cases to tumor-associated inflammation and fibrosis.77 However, rates of open resection and complications were favorably compared with surgery following NACT.77 Elsewhere, lung resection for residual disease following ICI for advanced malignancy was reported as feasible although potentially technically challenging, but with acceptable complication rates.78 An analysis of surgical outcomes from TOP1201 reported no increased adverse outcomes compared with those receiving NACT alone.79 In LCMC3, only 15 of 101 participants initially planned for minimally invasive surgery required conversion to thoracotomy, with surgery following NAICI deemed feasible.46 In a phase II trial of neoadjuvant atezolizumab plus chemotherapy, 46% resections were performed via video-assisted thoracoscopic surgery with no complications attributable to neoadjuvant treatment.44

FUTURE DIRECTIONS

Neoadjuvant therapy for cancer provides a range of potential benefits, including the ability to assess on-treatment response, reduce the tumor bulk prior to surgery, and enhance tolerability. Moreover, the nature of immunotherapy provides additional theoretical rationale for neoadjuvant administration. First, given that immune checkpoint blockade enhances T-cell activation on antigen encounter, administration of ICI in the neoadjuvant setting while the primary tumor remains in situ may result in a greater breadth of T-cell responses than in the adjuvant setting after surgery.80 81 Notably, the emergence of peripheral T-cell clones that were rarely detected in primary tumor was reported among patients receiving neoadjuvant nivolumab in NSCLC.82 Such expansion of ‘subdominant’ clones might reflect epitope-spreading arising from denovo priming after ICI administration, and could represent a new T-cell repertoire more ideally poised to contribute to effective antitumor immunity.82–84 Removal of the primary tumor also likely removes the risk of T-cell exhaustion resulting from chronic antigenic exposure. In support of this, preclinical studies discussed above have shown that NAICI induces a significantly higher level of tumor antigen-specific circulating T-cells compared with the corresponding adjuvant ICI, resulting in a long-lasting effector-memory T cell pool. The long-term persistence of tumor antigen-specific clonal T-cells seen in animal models of effective neoadjuvant therapy, and in patients who derived durable benefit from immune checkpoint blockade in NSCLC and colorectal cancer, suggest that tumor-specific T-cell memory might be a consequence.28 37 85

Unanswered questions relate to the ideal timing and duration of immunotherapy administration, including...
number of preoperative doses and scheduling with respect to surgery, predictive biomarkers to direct selection of therapy (eg, monotherapy ICI vs combination chemoinmunotherapy), and the requirement for adjuvant or ‘consolidation’ therapies (either chemotherapy and/or radiotherapy). Protocols to date generally recommend the standard administration of adjuvant chemotherapy or RT according to usual institutional practice following NAICI in NSCLC trials, where these have not already been given preoperatively. A concern about chemotherapy following NAICI could relate to the traditional understanding of the immunosuppressive and lymphodepleting nature of cytotoxic chemotherapy, although the interplay between chemotherapy and the immune system is nuanced. One preclinical study failed to demonstrate benefit of adjuvant ICI after NAICI. The PRADO expansion cohort of the OpACIN-neo trial (NCT02977052), where adjuvant ICI will be withheld in patients sustaining excellent response to NAICI, may provide insight into this question in the setting of resectable stage 3 melanoma. Given that ICI conveys risks of irAE that can prove life-threatening or permanent, the identification of robust biomarkers to determine who will benefit from therapy remains a priority. As attempts are made to intensify neoadjuvant treatment, such as through ICI-ICI combinations or ICI combined with other therapies, emergence of significant AEs may be a concern. Thorough reports of safety endpoints will be crucial in the widespread acceptance of NAICI. Importantly, validating MPR as a surrogate endpoint for survival would streamline efficacy assessment and accelerate the evaluation of novel neoadjuvant approaches.

Looking forward, an area for improvement in the trial design of NAICI studies in NSCLC could relate to greater collaboration between investigators, as has been achieved in melanoma research with the International Neoadjuvant Melanoma Consortium. Currently, there is marked heterogeneity in trial design and endpoints across phase 2 and 3 studies, and improved collaboration to enable reasonable comparability across trials, including in the accompanying translational research, would benefit to the field.

**SEARCH STRATEGY**

We searched ClinicalTrials.gov for all studies in all countries relating to ‘NSCLC’ (condition or disease) and ‘neoadjuvant’ (other term) with further filters applied: recruiting (status); phase 3 (study phase). Individual trial records were then reviewed to select those employing immunotherapy in resectable disease.

**Author affiliations**

1School of Clinical Sciences, Monash University Faculty of Medicine Nursing and Health Sciences, Clayton, Victoria, Australia  
2Medical Oncology, Monash Health, Clayton, Victoria, Australia  
3Immunology in Cancer and Infection, QIMR Berghofer Medical Research Institute, Herston, Queensland, Australia  
4Medical Oncology, Peter MacCallum Cancer Centre, Melbourne, Victoria, Australia  
5Medical Oncology, Westmead Hospital, Westmead, New South Wales, Australia  
6Faculty of Medicine, The University of Sydney, Sydney, New South Wales, Australia  
7Medical Oncology, Royal North Shore Hospital Northern Sydney Cancer Centre, St. Leonards, New South Wales, Australia  
8Medical Oncology, Princess Alexandra Hospital, Woolloongabba, Queensland, Australia  
9School of Biomedical Sciences, Queensland University of Technology, Brisbane, Queensland, Australia  
10Medical Oncology, Royal Brisbane and Women’s Hospital, Herston, Queensland, Australia  
11Faculty of Medicine, The University of Queensland, Herston, Queensland, Australia  
12Medical Oncology, The Prince Charles Hospital, Chermside, Queensland, Australia

**Contributors** Substantial contribution to conception and design of work: all authors. Drafting the work and/or revising it critically: all authors. Final approval of the version to be published: all authors. Agreement to be accountable for all aspects of the work: all authors.

**Funding** The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

**Competing interests** BGMR: research funding (institution): Amgen. Advisory board or consultation: Roche, AstraZeneca, Bristol-Myers Squibb, Merck Sharpe and Dohme, Pfizer, Eisai, Takeda, Boehringer-Ingelheim. BJS: Advisory board/honoraria: Roche/Pfizer/Geneintech, AstraZeneca, Pfizer, Novartis, Bristol-Myers Squibb, Merck, Amgen, Lilly oncology. EA: research funding (institution): Amgen. Non-financial support: Amgen, Bristol-Myers Squibb. KG: Advisory board/honoraria: Pfizer, Roche, AstraZeneca, Boehringer-Ingelheim, Bristol Myers Squibb, Merck, Lilly Oncology, Novartis, Janssen, Yuhan, Teva, Amgen and Naterra; shareholder and board member: Carpe Pharmaceuticales; Carpe Vita Pharmaceuticals; DGC Diagnostics. NP: research funding (institution): Bayer. Pfizer. Advisory board or consultation: Bristol-Myers Squibb, MSD, Merck-KGaA, Boehringer-Ingelheim, AstraZeneca, Roche, Bayer, Novartis, Merck-Serono, Pfizer, Takeda, Iopen. RH: Advisory board: AstraZeneca, Bristol-Myers Squibb, Eli Lilly, Merck, Merck Sharpe and Dohme, Novartis, Inc, Pfizer, Roche, Segen; Speaker honorarium: Merck Sharpe and Dohme, Novartis, Roche.

**Patient consent for publication** Not required.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Open access** This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See http://creativecommons.org/licenses/by-nc/4.0/.

**ORCID iD**

Elizabeth Ahern http://orcid.org/0000-0002-2062-5695

**REFERENCES**

Open access


57 Second affiliated Hospital, S. O. M. Z. U. (2022).


60 University Hospital, E. (2023).


66 Sidney Kimmel Comprehensive Cancer Center at Johns, H. & AstraZeneca 2021.


<table>
<thead>
<tr>
<th>Trial name/identifier</th>
<th>Trial design</th>
<th>Neoadjuvant trial intervention</th>
<th>Primary endpoint(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NEOMUN, NCT03197467</td>
<td>Phase 2, single-arm, n=30 Stage 2-3A</td>
<td>Pembrolizumab (2 cycles)</td>
<td>Safety, feasibility, tumour response (RECIST, PET and Junker criteria)</td>
</tr>
<tr>
<td>PRINCEPS, NCT02994576</td>
<td>Phase 2, single-arm, n=60 Stage 1B-3A (non-N2)</td>
<td>Atezolizumab (1 cycle)</td>
<td>Rate free of major toxicities or morbidities</td>
</tr>
<tr>
<td>NCT03732664</td>
<td>Phase 1, single-arm, n=40 Stage 1A-3A</td>
<td>Nivolumab (3 cycles)</td>
<td>Safety and AE rate</td>
</tr>
<tr>
<td>IoNESCO, NCT03030131</td>
<td>Phase 2, single-arm, n=81 Stage 1B-2B</td>
<td>Durvalumab (3 cycles)</td>
<td>R0 resection rate</td>
</tr>
<tr>
<td>TOP 1501, NCT02818920</td>
<td>Phase 2, single-arm, n=35 Stage 1B-3A</td>
<td>Pembrolizumab (2 cycles) + adjuvant pembrolizumab</td>
<td>Surgical feasibility rate</td>
</tr>
<tr>
<td>NEOpredict, NCT04205552</td>
<td>Phase 2, two-arm, randomized, open-label, n=60 Stage 1-3A</td>
<td>Nivolumab ± relatlimab (2 cycles)</td>
<td>Feasibility</td>
</tr>
<tr>
<td>POPCORN, ACTRN</td>
<td>Phase 1B/2, two-arm,</td>
<td>Nivolumab ± denosumab (2 cycles)</td>
<td>Pharmacodynamic correlates of</td>
</tr>
<tr>
<td>Trial ID</td>
<td>Study Design</td>
<td>Treatment Description</td>
<td>Outcome Measures</td>
</tr>
<tr>
<td>------------------</td>
<td>-------------------------------</td>
<td>---------------------------------------------------------------------------------------</td>
<td>------------------</td>
</tr>
<tr>
<td>12618001121257</td>
<td>Randomized, open-label, n=30</td>
<td>Stage 1-3A neoadjuvant therapy</td>
<td>NADIM-II</td>
</tr>
<tr>
<td>NCT03838159</td>
<td>Phase 2, two-arm, randomized, open-label, n=90</td>
<td>Potentially resectable stage 3A-B Platinum-doublet NACT ± neoadjuvant/adjuvant nivolumab</td>
<td>pCR</td>
</tr>
<tr>
<td>NCT03081689</td>
<td>Phase 2, single-arm, n=46</td>
<td>Stage 3A (N2) Platinum-doublet NACT + neoadjuvant/adjuvant nivolumab (1 neoadjuvant cycle)</td>
<td>PFS</td>
</tr>
<tr>
<td>NCT02572843</td>
<td>Phase 2, single-arm, n=68</td>
<td>Stage 3A (N2) Platinum-doublet NACT + neoadjuvant/adjuvant durvalumab (2 neoadjuvant cycles)</td>
<td>EFS</td>
</tr>
<tr>
<td>NCT03237377</td>
<td>Phase 2, single-arm, n=32</td>
<td>Stage 3A Durvalumab + thoracic RT (2 cycles, 45Gy) Possibility of tremelimumab expansion arm</td>
<td>Toxicity, feasibility</td>
</tr>
<tr>
<td>NCT03871153</td>
<td>Phase 2, single-arm, n=25</td>
<td>Stage 3 (N2) Chemoimmunoradio-therapy (platinum doublet + durvalumab + thoracic RT) (3 cycles, 45-61.2Gy) + adjuvant durvalumab</td>
<td>pCR</td>
</tr>
</tbody>
</table>

Supplementary Table 1. Ongoing phase 1-2 trials employing neoadjuvant immunotherapy in non-small cell lung cancer. NACT (neoadjuvant chemotherapy), RT
(radiotherapy), Gy (Gray), RECIST (Response Evaluation Criteria in Solid Tumors), PET (positron-emission tomography), AE (adverse events), pCR (pathological complete response), PFS (progression-free survival), EFS (event-free survival).
<table>
<thead>
<tr>
<th>Trial name/identifier</th>
<th>Trial design</th>
<th>Adjuvant trial intervention</th>
<th>Primary endpoint(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BR.31, NCT02273375</td>
<td>Phase 3, two-arm, randomized, double-blinded, n=1,360</td>
<td>Durvalumab vs placebo for up to 12 months in addition to standard-of-care adjuvant chemotherapy and/or radiotherapy in patients with resected stage 1B (primary $\geq$ 4cm) – 3A NSCLC</td>
<td>DFS in PD-L1 positive patients, DFS in all patients</td>
</tr>
<tr>
<td>PEARLS (KEYNOTE-091), NCT02504372</td>
<td>Phase 3, two-arm, randomized, double-blinded, n=1,080</td>
<td>Pembrolizumab vs placebo for up to 12 months in patients with resected stage 1B (primary $\geq$ 4cm) – 3A NSCLC</td>
<td>DFS</td>
</tr>
<tr>
<td>IMpower010, NCT02486718</td>
<td>Phase 3, two-arm, randomized, open-label, n=1,280</td>
<td>Atezolizumab in intervention group for up to 12 months following completion of adjuvant cisplatin-based chemotherapy in patients with resected stage 1B (primary $\geq$ 4cm) – 3A NSCLC</td>
<td>DFS in PD-L1-selected populations within various subgroups (stage 2-3A subpopulation, stage 2-3A participants, and ITT population)</td>
</tr>
<tr>
<td>ANVIL, NCT02595944</td>
<td>Phase 3, two-arm, randomized, open-label, n=903</td>
<td>Nivolumab in intervention group for up to 12 months following completion of any adjuvant chemotherapy or radiotherapy in patients with resected stage 1B</td>
<td>DFS and OS, each stratified by PD-L1 expression</td>
</tr>
</tbody>
</table>
(primary ≥ 4cm) – 3A NSCLC not harbouring EGFR mutations (exon 19 deletion or exon 21 L858R) or ALK rearrangement

| **Supplementary Table 2.** Phase 3 trials of adjuvant immunotherapy in resected non-small cell lung cancer (NSCLC). Note: in all trials, UICC version 7 staging was used, where stage 3A would now correlate with 3B in the case of pT3N2M0 disease, although these patients were still included. DFS, disease-free survival; PD-L1, programmed death-1-ligand 1; OS, overall survival; EGFR, epithelial growth factor receptor; ALK, anaplastic lymphoma kinase. |