

Treatment Options for Advanced Non-Small Cell Lung Cancer: Effectiveness and Value

Draft Evidence Report

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Institute for Clinical and Economic Review



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About ICER

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The Midwest Comparative Effectiveness Public Advisory Council (Midwest CEPAC) – a core program of ICER – provides a public venue in which the evidence on the effectiveness and value of health care services can be discussed with the input of all stakeholders. Midwest CEPAC seeks to help patients, clinicians, insurers, and policymakers interpret and use evidence to improve the quality and value of health care.

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List of Acronyms Used in this Report

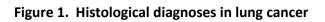
AE	Adverse event
AJCC	American Joint Committee on Cancer Staging
ASCO	American Society of Clinical Oncology
BI	Budget impact
BSA	Body surface area
CI	Confidence interval
CIS-PEM	Cisplatin+pemetrexed
COPD	Chronic obstructive pulmonary disease
ECOG PS	Eastern Cooperative Oncology Group Performance Status score
EGFR	Epidermal growth factor receptor
FDA	US Food and Drug Administration
HR	Hazard ratio
HrQoL	Health-related quality of life
ICER	Incremental cost-effectiveness ratio
irAE	Immune-related adverse events
ITT	Intention to treat
NCCN	National Comprehensive Cancer Network
NICE	National Institute for Health and Care Excellence
NMA	Network meta-analysis
NSCLC	Non-small cell lung cancer
OR	Odds ratio
ORR	Objective response rate
OS	Overall survival
PD-1	Programmed death 1 receptor
PD-L1	Programmed death ligand 1
PF	Progression-free
PFS	Progression-free survival
PRISMA	Preferred reporting items for systematic reviews and meta-analyses
PRO	Progression
PSA	Probabilistic sensitivity analysis
QALY	Quality-adjusted life-year
QoL	Quality of life
RCT	Randomized controlled trial
RECIST	Response evaluation criteria in solid tumors
TEAE	Treatment-emergent adverse event
ТКІ	Tyrosine kinase inhibitor
TNM -	Tumor, lymph nodes, metastasis
Tx	Treatment
USPSTF	US Preventive Services Task Force
WAC	Wholesale acquisition price

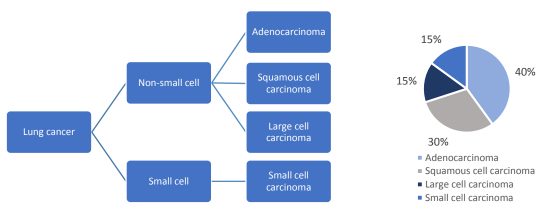
1. Background

Introduction

Lung cancer is the number one cause of cancer death in the United States, expected to cause 158,000 deaths in 2016, and accounting for 26.5% of all cancer deaths.¹ It is the second most common cancer in both men (after prostate cancer) and women (after breast cancer), with 118,000 and 106,000 new cases expected in 2016, respectively.² The median age at death is 72.¹ Lung cancer rates reflect smoking behavior, and the incidence of lung cancer peaked in men in 1992 with 69.5 cases per 100,000 and in women in 2005 with 53.8 cases per 100,000; those rates declined by 2013 to 52.2 and 47.7 cases per 100,000, respectively, reflecting earlier declines in the prevalence of smoking.^{3,4}

Lung cancer includes different pathological types (see Figure 1), broadly divided into small-cell lung cancer and non-small cell lung cancer (NSCLC).⁵ NSCLC makes up about 85% of lung cancers and comprises mainly squamous cell carcinoma, adenocarcinoma, and large-cell lung cancer.⁶ Stage at diagnosis is a primary factor in patient survival, and patients with NSCLC commonly present with advanced disease (i.e., distant spread, malignant effusion, or bilateral lung disease); 24% have regional spread at presentation, and 55% have distant spread.³ Prognosis is generally poor at diagnosis; five-year survival from 2006-2012 was 31.4% in patients with regional spread and 4.9% in patients with distant spread.





Source: American Cancer Society. Lung Cancer (Non-Small Cell). 2016; <u>http://www.cancer.org/acs/groups/cid/documents/webcontent/003115-pdf.pdf</u>. Accessed August 15, 2016.

Previously, advanced NSCLC was treated with chemotherapy with a platinum-based chemotherapy doublet (e.g., cisplatin+paclitaxel, carboplatin+gemcitabine, etc.). In recent years, the treatment of

some advanced NSCLCs has changed based on the determination of driver mutations in tumors. Among the most common of these driver mutations that affect therapeutic decisions are those involving the kinase region of the epidermal growth factor receptor (EGFR). More recently, immunotherapy aimed at altering checkpoint inhibition through the programmed death 1 (PD-1) receptor or its ligand (PD-L1) shows promise in at least some patients with NSCLC. Questions remain, however, regarding the appropriate sequence of treatment with these newer agents, the role of certain tests to inform treatment decisions, and management of the costs of these therapies.

Scope of the Assessment

This assessment evaluates the health and economic outcomes of certain tyrosine kinase inhibitors (TKIs) and programmed death 1 (PD-1) agents in the treatment of advanced non-small cell lung cancer (NSCLC). The effects of both classes of agents are evaluated in EGFR-positive (EGFR+) NSCLC, and PD-1 agents are evaluated in NSCLC without a driver mutation (EGFR-). The scope is described below using the PICOTS (Population, Intervention, Comparators, Outcomes, Timing, and Settings) framework.

Analytic Framework

The analytic framework for this assessment is depicted in Figure 2.

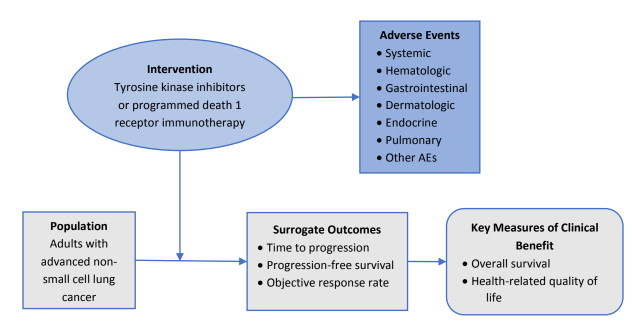


Figure 2. Analytic Framework: Management of Advanced Non-Small Cell Lung Cancer

Populations

The four populations of focus for the review were adults with advanced NSCLC who:

- P1) Have an EGFR+ tumor and have not previously been treated for advanced disease
- P2) Have a tumor without a driver mutation and have not previously been treated for advanced disease
- P3) Have a tumor without a driver mutation that has progressed after first-line treatment with a platinum-based chemotherapy doublet (e.g., cisplatin+paclitaxel, carboplatin+gemcitabine, etc.)
- P4) Have an EGFR+ tumor that has progressed after first-line or first- and second-line TKI therapy (patients who do not develop a T790M mutation will only receive first-line TKI therapy)

Interventions

- P1) The TKIs erlotinib, gefitinib, and afatinib
- P2) A treatment sequence of PD-1 immunotherapy (i.e., nivolumab, pembrolizumab, or atezolizumab), followed by a platinum-based chemotherapy doublet at the time of progression
- P3) PD-1 immunotherapy (after progression on a platinum-based chemotherapy doublet)
- P4) PD-1 immunotherapy (after progression on first-line or first- and second-line TKI therapy)

Comparators

- P1) A platinum-based chemotherapy doublet
- P2) A treatment sequence of a platinum-based chemotherapy doublet, followed by PD-1 immunotherapy at the time of progression
- P3) Single-agent chemotherapy (e.g., docetaxel)
- P4) A platinum-based chemotherapy doublet

<u>Outcomes</u>

This review examined key clinical outcomes that occur in all four populations of patients being treated for advanced NSCLC, including surrogate outcomes common to cancer trials. Outcomes of interest included:

- Overall survival
- Disease progression-related measures (progression-free survival, time to progression)
- Objective response rate
- Symptom control
- Health-related quality of life
- Treatment-related adverse events

- Rates of key adverse events by type (e.g., systemic, gastrointestinal, dermatologic, etc.)
- o Rates of Grade 3 or 4 adverse events
- Discontinuation due to adverse events
- o Treatment-related deaths

<u>Timing</u>

Evidence on intervention effectiveness and harms was derived from studies of any duration.

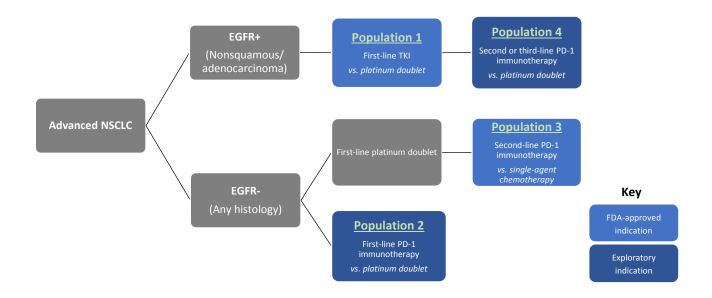
<u>Settings</u>

All relevant settings were considered, including inpatient, clinic, and office settings.

2. The Topic in Context

As discussed above, the prognosis in patients with advanced NSCLC has been poor, and new therapies are needed for this common malignancy. Although chemotherapy can extend survival, it is not curative in patients with advanced disease, and many patients may be unable to tolerate the side effects of the most potent regimens. These potent chemotherapy regimens are typically platinum-based chemotherapy doublets (e.g., cisplatin + paclitaxel, carboplatin + gemcitabine, etc.), and they continue to be recommended as first-line therapy for patients with advanced NSCLC without a driver mutation.⁷ Figure 3 presents the populations and therapies of interest for this review.

Figure 3. Populations and therapies of focus



Tyrosine Kinase Inhibitors (TKIs)

Mutations affecting the tyrosine kinase region of the epidermal growth factor receptor (EGFR) are found in approximately 10 to 15% of patients with adenocarcinoma in the United States, but in up to 50% of patients from Asia.^{5,8} EGFR mutations are more common in non-smokers with NSCLC and less common in squamous cell NSCLC (approximately 2.7% with EGFR mutations).⁷

For NSCLC patients with EGFR mutations (referred to as EGFR+ NSCLC), major guidelines, such as those from the National Comprehensive Cancer Network (NCCN), list tyrosine kinase inhibitors (TKIs) as first-line therapy.⁷ The main TKIs used as first-line therapy for advanced NSCLC include

afatinib (Gilotrif[®], Boehringer Ingelheim), erlotinib (Tarceva[®], Genentech), and gefitinib (Iressa[®], AstraZeneca). The US Food and Drug Administration (FDA) approved both erlotinib and afatinib for use in the first-line setting in 2013, and granted approval to gefitinib in 2015; these medications are approved for treatment of advanced NSCLC with the two most common EGFR mutations: frame deletions in exon 19 and the substitution of arginine for leucine at codon 858 in exon 21 (L858R).⁸ There is some evidence that the type of EGFR mutation may influence response to TKI therapy, including possible differences in response between exon 19 and L858R mutations.⁸⁻¹⁰

TKIs are administered orally once daily (see Table 1), generally until disease progression. The most common adverse reactions are rash, which can be severe, and diarrhea.^{11,12} In general, however, rates of serious adverse events are much lower with TKI therapy than with a platinum doublet. A course of treatment with first-line TKI therapy typically costs approximately \$90,000 per year.¹³

Platinum-based chemotherapy doublets have been the comparators for most trials of TKIs.¹⁰ There are few head-to-head trials comparing TKIs.

As discussed in detail in section 4, trials of TKIs have generally shown improvements in progressionfree survival (PFS) compared with chemotherapy, but no improvement in OS, which is likely due to the high rates of crossover between treatment arms that occurred in these trials.

Despite treatment with a TKI, nearly all patients with advanced NSCLC will eventually progress.⁵ A common mechanism of TKI resistance is a T790M mutation. Commonly, patients who progress and have a T790M mutation are treated second-line with osimertinib (Tagrisso[®], AstraZeneca), a TKI that is effective in EGFR+ tumors with this mutation.⁷ For patients who progress on first-line TKI therapy but do not have a T790M mutation, or who progress on osimertinib, guidelines suggest proceeding with chemotherapy doublet treatment as in patients without a driver mutation.⁷

Table 1. TKIs of interest for the evidence review

	Administration	Recommended dose	Treatment Duration	Unit Price (USD) ^a
Afatinib (Gilotrif®) Boehringer Ingelheim	Oral	40 mg, once daily	Until progression or no longer tolerated by the patient	\$233.05/tablet
Erlotinib (Tarceva®) Genentech	Oral	150 mg, once daily	Until progression or unacceptable toxicity	\$241.52/tablet
Gefitinib (Iressa®) AstraZeneca	Oral	250 mg, once daily	Until progression or unacceptable toxicity	\$241.20/tablet

 α Unit price reflects the wholesale acquisition price listed on Red Book Online (Greenwood Village, CO: Truven Health Analytics. http://www.micromedexsolutions.com/. Accessed July 19, 2016).

PD-1 immunotherapy

Tumor cells can produce substances that alter the immune response to the tumor, such as by affecting a regulatory "checkpoint" or brake on the T cell response to the tumor, and thus allowing the tumor to evade the immune system. Immunotherapy aimed at inhibiting such a checkpoint through the PD-1 receptor or its ligand, PD-L1, shows promise in at least some patients with NSCLC.¹⁴ Agents focused on this pathway include nivolumab (Opdivo®, Bristol-Myers Squibb) and pembrolizumab (Keytruda®, Merck), which are antibodies to PD-1, as well as atezolizumab (Tecentriq®, Genentech), an antibody to PD-L1. We use the term "PD-1 immunotherapy" to refer to both groups of antibodies. Both nivolumab and pembrolizumab received FDA approval for NSCLC in 2015. Atezolizumab, which was approved for advanced or metastatic urothelial carcinoma in May of 2016, is not yet indicated for NSCLC; the FDA is expected to issue a final decision on the use of atezolizumab in NSCLC by October 19, 2016. PD-1 immunotherapy is recommended as second-line treatment in patients with advanced NSCLC without a driver mutation who progress on a chemotherapy doublet.⁷

A minority of patients respond to PD-1 immunotherapy, but a substantial proportion of those who do respond appear to have prolonged responses and improved survival.^{14,15} Tumor expression of PD-L1 appears to be helpful in selecting appropriate patients for PD-1 based therapies; however, there are concerns about the comparability of various methods used to assess levels of expression.¹⁴ The FDA-labeled indication for pembrolizumab is for patients whose tumors express PD-L1. While nivolumab does not carry such a restriction, response rates are higher in tumors with PD-L1 expression with nivolumab as well.¹⁴ However, some tumors that do not express PD-L1 respond to PD-1 immunotherapy such that it is likely that only treating patients based on PD-L1 expression would result in missing an opportunity for therapy in a percentage of patients.

Pembrolizumab is administered every three weeks, and atezolizumab is likely to be approved using an every 3-week dosing schedule; nivolumab is administered every two weeks (see Table 2). Both are given by intravenous infusion until disease progression. Fatigue is common with these agents, and more serious immune-mediated events, including pneumonitis and encephalitis, have been seen. These immune events are uncommon, and serious adverse events overall are much less common with PD-1 immunotherapy than with docetaxel.

Most patients studied in trials of PD-1 immunotherapy have received prior treatment with a chemotherapy doublet, whether or not they were EGFR+ and/or had received prior TKI therapy.¹⁶⁻¹⁸ The alternative treatment in this setting would typically be single-agent chemotherapy with an agent that was not used in the original doublet, such as docetaxel.

Recently, researchers and clinicians have begun exploring the use of PD-1 agents in patients who have not received a chemotherapy doublet.¹⁹ This includes using PD-1 immunotherapy as first-line treatment in patients with NSCLC without a driver mutation or as third-line therapy (after osimertinib) in patients with EGFR+ NSCLC.

A course of PD-1 immunotherapy has been estimated to cost approximately \$150,000 per year.¹³ In addition to questions of the comparative effectiveness of these agents, both among the agents and compared with alternative therapies, the use of PD-L1 levels to select patients for treatment is likely to affect estimates of cost-effectiveness.²⁰

Multiple trials are currently underway looking at PD-1 immunotherapy in advanced NSCLC. Issues being studied include the use of these therapies first line, and the use of PD-1 immunotherapies in combination with each other or with other therapies. We summarize relevant ongoing studies in Appendix H.

	Administration	Recommended dose	Treatment Duration	Price (USD) ^β
Atezolizumab (Tecentriq®) <i>Genentech</i>	Intravenous infusion	1200 mg on first day of every 3-week cycle $^{\alpha}$	Until disease progression or unacceptable toxicity	• \$7.18/mg • \$8,620.00/20 mL vial
Nivolumab (Opdivo®) Bristol-Myers Squibb	Intravenous infusion	3 mg/kg over 60 minutes every 2 weeks	Until disease progression or unacceptable toxicity	• \$24.70/mg • \$2,470.00/10 mL vial • \$988.19/4 mL vial
Pembrolizumab (Keytruda®) <i>Merck</i>	Intravenous infusion	2 mg/kg over 30 minutes every 3 weeks	Until disease progression or unacceptable toxicity	• \$43.81/mg • \$4,380.74/4 mL vial

Table 2. PD-1 immunotherapies of interest for the evidence review

 α Atezolizumab has not yet been approved for NSCLC in the US. Dose represents that used in POPLAR trial²¹ (as well as approved dose for urothelial cancer); β Price reflects the wholesale acquisition price listed on Red Book Online (Greenwood Village, CO: Truven Health Analytics. http://www.micromedexsolutions.com/. Accessed July 19, 2016).

Definitions

We provide the following definitions to help with interpretation of the study results presented throughout this report.

Response Criteria: Note, trials prior to 2009 typically used response evaluation criteria in solid tumors (RECIST) version 1.0 criteria,²² while trials after 2009 typically used RECIST version 1.1 criteria.²³ The definitions below reflect version 1.1; for most cases of patients with advanced NSCLC, there is close agreement between the two versions in assessing measures of response.²⁴

- *Progressive disease:* At least a 20% increase in the sum of diameters of target lesions, taking as reference the smallest sum on study (this includes the baseline sum if that is the smallest on study); in addition to the relative increase of 20%, the sum must also demonstrate an absolute increase of at least 5 mm **OR** unequivocal progression of existing non-target lesions **OR** the appearance of one or more new lesions.
- *Complete response:* Disappearance of all target and non-target lesions, and normalization of tumor marker levels. Any pathologic lymph nodes (whether target or non-target) must have a reduction in the short axis to <10 mm.
- *Partial response:* At least a 30% decrease in the sum of the diameters of target lesions and not meeting criteria for progressive disease or complete response.
- *Objective response:* Complete response or partial response.
- Advanced disease: Stage IIIB or IV lung cancer according to American Joint Committee on Cancer Staging (AJCC) TNM staging (editions 6 or 7).^{a25,26}
 - Stage IIIb NSCLC: Cancer is present in the lung and lymph nodes (on the opposite side of the chest from the affected lung or near the collarbone), has spread to different lobes of the same lung, or has grown into the structures surrounding the lung (i.e., the mediastinum, heart, aorta, trachea, esophagus, backbone, or carina).
 - *Stage IV NSCLC*: Cancer has spread to both lungs, to fluid in the area around the lungs or heart, or to another part of the body, such as the brain, bones, liver or adrenal glands.

Eastern Cooperative Oncology Group (ECOG) performance status: a measure of functional status and ability to perform activities of daily living on a 6-point scale: 0 (fully active); 1 (restricted only in strenuous activity); 2 (ambulatory and capable of self-care but unable to work); 3 (capable of only limited self-care, confined to bed or chair >50% of waking hours); 4 (completely disabled); 5 (dead).²⁷

^a Tumor, Lymph Node, Metastasis (TNM). The TNM system for lung cancer is based on the size of the original tumor (**T**) and whether it has grown into nearby areas, whether the cancer is present in the lymph nodes (**N**), and whether the cancer has spread to other organs (metastasis-**M**).

Toxicity criteria for adverse events: Grade 3 adverse events are severe or medically significant, but not life threatening. Hospitalization or prolongation of hospitalization is required, and the event limits the patient's ability for self-care. Grade 4 adverse events are life threatening and require urgent intervention.

Insights Gained from Discussions with Patients and Patient Groups

Lung cancer, unlike most other cancers and serious illnesses, is a disease where the patient often feels blamed for the illness because of the strong association with prior smoking behavior. As such, lung cancer has an unusual stigma that affects the interactions that patients with lung cancer have with family, friends, and providers. We were told that patients with lung cancer and breast cancer are at the highest risk for depression among cancer patients, and that lung cancer patients have higher levels of fatigue, distress, and anxiety than other cancer patients. High levels of anxiety may be due, in part, to breathing difficulties and other symptoms of the disease.

Because of prior smoking, lung cancer patients are at particularly high risk for comorbidities that can affect their ability to participate in clinical trials and/or receive specific therapies. Particularly important and common comorbidities include vascular disease (cardiovascular, peripheral arterial, and cerebrovascular) and chronic obstructive pulmonary disease (COPD). As such, the results of clinical trials in which these comorbidities are underrepresented may not generalize well to the patient population as a whole.

With TKI therapy in particular, there can be heightened anxiety around adverse events and reporting these events. Patients have heard that development of rash may be associated with better response to TKIs,²⁸ but also have concerns about therapy being discontinued if adverse events are reported. This may affect the frequency of adverse events reported in the published literature.

In certain settings, such as rural or low-income community clinics, patients may not receive the same care as they would in major medical centers. We heard that some patients do not receive molecular testing (or are not apprised of the results of such tests) and are being put directly on chemotherapy without regard to the individual clinical characteristics of their disease. Access to innovative and emerging therapies is out of reach for those who live far from major centers without the resources to travel for treatment. Such patients may be less likely to receive adequate support, participate in clinical trials where emerging treatment may be available, or receive education on their diagnosis, prognosis, and treatment options.

Multiple patient groups raised the issue of the "financial toxicity" of treatment for lung cancer, with patients and their families at high risk for suffering economic hardship or even bankruptcy. The financial toxicity of cancer reflects the cost of the medications, administration of the medications, travel to receive treatments and other care, missing work, and even requirements for new clothing

as a result of weight loss. As patients live longer, treatment costs accumulate over time. Additionally, as smoking has become less common in patients of higher socioeconomic status, lung cancer is now disproportionately affecting patients who have fewer resources available to deal with the illness. We heard that some of these patients just expect to die and so do not seek treatment at all.

Studies have not adequately addressed high rates of distress in NSCLC patients, and patients in clinical trials (who typically have significant resources provided by the trial and do not need to pay for medications) may not show similar levels of distress compared to patients treated outside of such trials.

Several groups commented that there is inadequate evidence from clinical trials on appropriate sequencing of therapies. We heard that as better therapies for NSCLC have been developed, patients have a goal of staying stable until the next trial/therapy becomes available. This is similar to the situation with HIV in the early 1990s.

3. Summary of Coverage Policies and Clinical Guidelines

All of the drugs under review in this report are covered by private insurers for use within their FDA labeled indications. The FDA labeled indication for pembrolizumab includes a companion diagnostic test to provide evidence of the expression of the protein PD-L1, which is required by payers. Some payers, such as Anthem, have developed treatment "pathways," or recommended regimens for providers. Anthem's pathway recommends afatinib or erlotinib for EGFR + first-line treatment, and nivolumab as second-line treatment for both squamous and non-squamous metastatic disease. We have also summarized here the clinical guidelines available for the treatment of advanced non-small cell lung cancer. We reviewed the National Comprehensive Cancer Network's (NCCN) guidelines for non-small cell lung cancer, version 4.2016, for treatments within scope. We also reviewed the most recent American College of Chest Physician (ACCP) guidelines, published in 2013.

National Comprehensive Cancer Network (NCCN)

Population P1: For patients with EGFR mutations, not previously treated, NCCN guidelines recommend the TKIs erlotinib, gefitinib, and afatinib for first line therapy.

Population P2: For patients without a driver mutation, not previously treated, NCCN guidelines recommend treating with a platinum-based chemotherapy doublet as first line therapy.

Population P3: For patients without a driver mutation who progress on a chemotherapy doublet, NCCN guidelines recommend second-line treatment with PD-1 immunotherapy, which currently includes nivolumab and pembrolizumab.

Population P4: For patients with EGFR mutations who have progressed after first-line or first- and second-line TKI therapy, NCCN guidelines recommend treating with a platinum-based chemotherapy doublet.

American College of Chest Physicians (ACCP)

(http://journal.publications.chestnet.org/article.aspx?articleID=1685102)

The American College of Chest Physicians published guidelines on the treatment of NSCLC in 2013. As a result, these guidelines do not include afatinib or any of the PD-1 immunotherapies, which had not yet been approved by the FDA. We included these guidelines as added context to the clinical environment. ACCP recommends, as a general approach, that patients with good performance status (PS) and stage IV non-small cell lung cancer should receive a platinum-based chemotherapy regimen. For patients that are EGFR+, ACCP recommends gefitinib or erlotinib as first-line therapy. ACCP also suggests erlotinib as a maintenance therapy for those patients that do not experience disease progression after 4 cycles of platinum-based double agent chemotherapy.

4. Comparative Clinical Effectiveness

4.1 Overview

To inform our analysis of the comparative clinical effectiveness of TKIs and PD-1 immunotherapy in the treatment of advanced NSCLC, we abstracted evidence from available clinical studies of these agents, whether in published, unpublished, or in abstract form.

Therapies of interest included:

- 1. TKIs for chemotherapy-naïve patients with an EGFR+ tumor (population 1 as described in Section 1)
 - Afatinib
 - Gefitinib
 - Erlotinib
- 2. PD-1 immunotherapy for patients without an EGFR+ tumor who are either chemotherapynaïve or have progressed after first-line treatment with a platinum doublet, and patients who have an EGFR+ tumor that has progressed after first- or second-line treatment with a TKI (populations 2-4 as described in Section 1)
 - Atezolizumab
 - Nivolumab
 - Pembrolizumab

As described previously in the Background section, comparators of interest included 1) platinumbased chemotherapy doublets for the first- and third-line EGFR+ treatment populations (populations 1 and 4), 2) platinum-based chemotherapy doublets for the first -line EGFR- treatment population (population 2) and 3) single-agent chemotherapy for the EGFR- population that has progressed after treatment with a platinum doublet (population 3). Our review focused on clinical benefits (i.e., overall and progression-free survival, biochemical response, and health-related quality of life), and potential harms (drug-related adverse events). We focused attention on both descriptive and quantitative analyses of these outcomes, including direct comparisons available from the individual trials as well as indirect comparisons between the newer regimens.

4.2 Methods

Study Selection

We included evidence from randomized controlled trials (RCTs), comparative observational studies, and high-quality systematic reviews where available. We excluded single-arm studies, studies

without an active control arm, and studies from an early clinical development phase (i.e., phase I). In recognition of the rapidly evolving evidence base for NSCLC, we supplemented our review of published studies with data from conference proceedings, regulatory documents, information submitted by manufacturers, and other grey literature that met ICER standards for review (for more information, see (http://icerreview.org/methodology/icers-methods/icer-value-assessment-framework/grey-literature-policy/).

Data Sources and Searches

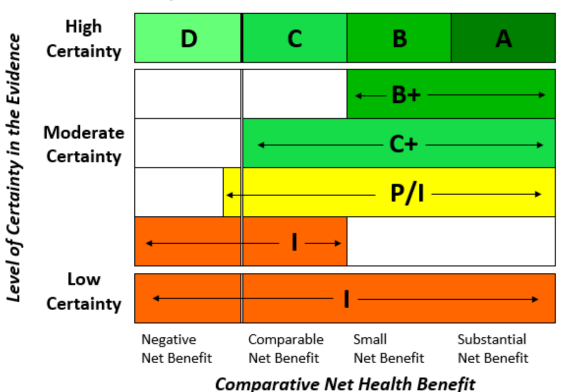
Procedures for the systematic literature review assessing the evidence on NSCLC regimens followed established best methods.²⁹ We conducted the review in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines.³⁰ The PRISMA guidelines include a checklist of 27 items, further detail of which is available in Appendix Table A1.

The timeframe for our search spanned the period from January 1996 to June 8, 2016 and focused on MEDLINE, EMBASE, and Cochrane-indexed articles. We limited each search to studies of human subjects and excluded articles indexed as guidelines, letters, editorials, narrative reviews, case reports, or news items. To supplement the above searches and ensure optimal and complete literature retrieval, we performed a manual check of the references of recent relevant reviews and meta-analyses. Further details on the search algorithm are available in Appendix Tables A2, A3, and A4. Additional searches were performed to identify relevant grey literature based on an organization and source checklist developed by the Canadian Agency for Drugs and Technologies in Health (https://www.cadth.ca/resources/finding-evidence/grey-matters). Other grey literature sources included sites deemed relevant specifically for NSCLC, such as clinical societies, research foundations, and advocacy organizations. Further information on methods for study selection, data extraction, quality assessment, assessment for publication bias, and our approach to meta-analyses of the data can be found in the appendices.

Assessment of Level of Certainty in Evidence

We used the <u>ICER Evidence Rating Matrix</u> (see Figure 4) to evaluate the evidence for a variety of outcomes. The evidence rating reflects a joint judgment of two critical components:

- The **magnitude** of the difference between a therapeutic agent and its comparator in "net health benefit" the balance between clinical benefits and risks and/or adverse effects AND
- The level of **certainty** in the best point estimate of net health benefit.³¹



Comparative Clinical Effectiveness

A = "Superior" - High certainty of a substantial (moderate-large) net health benefit

B = "Incremental" - High certainty of a small net health benefit

C = "Comparable"- High certainty of a comparable net health benefit

D = "Negative"- High certainty of an inferior net health benefit

B+ = "Incremental or Better" - Moderate certainty of a small or substantial net health benefit, with high certainty of at least a small net health benefit

C+ = "Comparable or Better" - Moderate certainty of a comparable, small, or substantial net health benefit, with high certainty of at least a comparable net health benefit

P/I = "Promising but Inconclusive" - Moderate certainty of a comparable, small, or substantial net health benefit, and a small (but nonzero) likelihood of a negative net health benefit

I = "Insufficient" - Either moderate certainty that the best point estimate of comparative net health benefit is comparable or inferior; or any situation in which the level of certainty in the evidence is low

4.3 Results

Study Selection

Our literature search identified 3,072 potentially relevant references (see Appendix Figure B1), of which 44 references met our inclusion criteria; these citations related to 17 individual studies and seven systematic reviews. Primary reasons for study exclusion included use of a combination regimen not approved by the FDA, comparison to a treatment that does not reflect current best practice (e.g., single-agent docetaxel in treatment-naïve EGFR+ patients), study population out of

our scope, and non-comparative study design. Overall, we identified 36 references relevant to population 1 (first-line EGFR+) and eight references germane to population 3 (second-line EGFR-); we did not identify any relevant references for populations 2 and 4 (first-line EGFR- and second- or third-line EGFR+, respectively). Details of the included studies are described in Appendix Table B1 and in the sections that follow; previous systematic reviews are described in Appendix G.

Tyrosine Kinase Inhibitors for First-Line Treatment of EGFR+ NSCLC

Our review of TKIs in patients with EGFR+ advanced NSCLC focused on first-line use of afatinib, erlotinib, and gefitinib (population 1 as described above). We assessed these therapies in relation to platinum-based chemotherapy doublets, which combined either cisplatin or carboplatin with gemcitabine, pemetrexed, paclitaxel, or docetaxel. The sections that follow discuss the overall survival (OS), progression-free survival (PFS), tumor response, symptom control, quality of life (QoL), and harms associated with the interventions of focus.

Summary

- Our review of the evidence on first-line TKI therapy for EGFR+ advanced NSCLC found inadequate evidence to distinguish between the three TKIs on patient-important outcomes such as OS and QoL.
- Evidence from RCTs indicates that all three agents provide statistically-significant improvements in PFS relative to platinum doublet chemotherapy. A head-to-head randomized trial, as well as our network meta-analysis, provide evidence that treatment with afatinib likely provides a small benefit in PFS compared with gefitinib; this is a surrogate endpoint and no statistically significant differences were seen in OS.
- RCTs comparing TKIs with platinum doublet chemotherapy had high rates of crossover, and showed no benefit in OS.
 - The most likely explanation for the lack of OS benefit seen in RCTs of TKIs with high rates of crossover is that treatment with a TKI improves survival *whenever* it is given in the sequence of therapy, such that patients randomized to chemotherapy who receive TKI therapy at the time of progression have similar survival in comparison to patients treated initially with a TKI.
 - Observational data suggest that first-line TKI therapy as a class *increases OS by approximately 8.9 months,* although there is substantial uncertainty in this figure.
 - Patients and clinicians should be aware that it is possible that the OS benefit with TKIs may be somewhat longer than this estimate in patients with tumors that have the exon 19 deletion, and somewhat shorter than this estimate in patients with tumors with the L858R mutation.

- Limitations in the evidence base preclude and definite conclusions as to whether the likely PFS benefit seen with afatinib over gefitinib would translate into a clinically important benefit in OS.
- Rash, diarrhea, and liver function abnormalities are the most common side effects with TKIs. All TKIs appear to be better tolerated than chemotherapy with platinum doublets, which have much higher rates of hematologic toxicity.
- QoL improvements were greater with TKI therapy than with chemotherapy.

For patients with EFGR+ advanced NSCLC, we have high certainty that TKI therapy provides at least a small net health benefit ("B+") relative to platinum chemotherapy.

Study Selection

Our literature search identified 36 references of afatinib, erlotinib, or gefitinib in chemotherapynaïve patients with an EGFR mutation; these citations related to 13 individual studies and five systematic reviews (see Appendix Table B1 and Appendix G). Ten of the 13 studies were rated fair quality and compared a TKI to a platinum doublet: two published Phase III RCTs of afatinib and four published Phase III trials each of erlotinib and gefitinib. An additional study evaluated gefitinib in comparison to standard chemotherapy, however this study was only published in a conference abstract and therefore not rated for quality. Finally, two studies (one good-quality phase IIb RCT and one fair-quality matched-pair case control study) directly compared two TKIs of interest (afatinib vs. gefitinib and gefitinib vs. erlotinib, respectively). Although the studies we deemed fair quality possessed many elements of a good quality study (e.g., comparability between study arms at baseline, use of valid instruments to evaluate outcomes, no differential attrition), we were concerned that the open-label design of these trials and high crossover rates could have potentially introduced biased estimates of treatment effect. We did not assign a quality rating to the remaining twelve documents, which were obtained from conference proceedings and previous systematic reviews. Appendix Table C1 presents the number of good, fair, poor, and unrated studies identified for each population of focus.

Key Studies

We considered 11 RCTs to be key studies of interest for this review, ten of which compared a TKI to a platinum-based chemotherapy doublet and one which compared afatinib to gefitinib (see Table 3). Important outcomes from each trial are also provided in Appendix Table B1, and described in further detail in the sections that follow. The trials specified similar inclusion criteria: each trial included treatment-naïve adult patients (≥18 years of age) with measurable, advanced (stage IIIb/IV) or recurrent NSCLC. All but three key trials³²⁻³⁴ were limited to EGFR+ patients; trials that did not restrict study inclusion criteria to EGFR+ patients provided results in this subgroup. The LUX-Lung 7 trial of afatinib vs. gefitinib provided the only direct randomized trial evidence comparing TKIs.

Trial populations were similar with respect to age, ECOG performance status, and disease stage: the median age was around 60 years, most patients had limited restrictions on their ability to perform daily activities (ECOG PS of 0-1), and around 90% of patients had stage IV NSCLC. With the exception of the TORCH trial, which compared erlotinib to cisplatin + gemcitabine, nearly all patients in each trial had a histological diagnosis of adenocarcinoma. The percentage of Asian patients was high in most trials, except for EURTAC and TORCH, which were conducted in European countries and Canada. Frequency of current or former smoking varied, and only one trial (First-SIGNAL trial of gefitinib vs. cisplatin + gemcitabine) was restricted to never-smokers. All of the key studies were open-label, and all but one trial (the phase IIb LUX-Lung 7 trial of afatinib vs. gefitinib) were phase III. There was no universal comparator treatment because standard of care for NSCLC varied by country, but cisplatin or carboplatin plus gemcitabine was the most commonly used comparator.

Table 3. Key studies: TKIs

Key Trials	Patient Characteristics	Treatment	Comparator	Harms (Treatment Arm)
LUX-Lung 3^{35,36α} Median f/u: 16.4 m	Median age: 61; Asian: 72% ECOG PS=1: 61% Never smoker: 68% Stage IV: 89%	Afatinib (n=230) Median OS: 28.2 m Median PFS: 11.1 m	Cisplatin+Pemetrexed (n=115) Median OS: 28.2 Median PFS: 6.9 m	D/C due to AEs: 10% TEAE ≥ Grade 3: 49% Tx-related deaths: 4
LUX-Lung 6^{36,37α} Median f/u: 16.6 m	Median age: 58; Asian: 100% ECOG PS=1: 76% Never smoker: 77% Stage IV: 94%	Afatinib (n=242) Median OS: 23.1 m Median PFS: 11.0 m	Cisplatin+Gemcitabine (n=122) Median OS: 23.5 m Median PFS: 5.6 m	D/C due to AEs: 9% TE-SAEs: 6% Tx-related deaths: 1
LUX-Lung 7³⁸ Median f/u: 27.3 m	Median age: 63; Asian: 57% ECOG PS=1: 69% Never smoker: 67% Stage IV: 97%	Afatinib (n=160) Median OS: 27.9 m Median PFS: 11.0 m	Gefitinib (n=159) Median OS: 25.0 m Median PFS: 10.9 m	D/C due to AEs: 11% TE-SAEs: 11% Tx-related deaths: 0
IPASS^{33,39} Median f/u: 17.0 m	Median age: 57; Asian: 100% ECOG PS=1: 64% Never smoker: 94% Stage IV: 76%	Gefitinib (n=132) ^β Median OS: 21.6 m Median PFS: 9.5 m	Carboplatin+Paclitaxel (n=129) ^β Median OS: 21.9 m Median PFS: 6.3 m	D/C due to AEs: 7% SAEs: 16% AE-related deaths: 4%
NEJ002^{40,41} Median f/u: 23.1 m	Mean age: 63; Asian: 100% ECOG PS=1: 50% Never smoker: 62% Stage IV: 75%	Gefitinib (n=114) Median OS: 30.5 m Median PFS: 10.8 m	Carboplatin+Paclitaxel (n=114) Median OS: 23.6 m Median PFS: 5.4 m	D/C due to AEs: NR AEs ≥ Grade 3: 41% Tx-related deaths: NR
WJTOG3405^{42,43} Median f/u: 59.1 m	Median age: 64; Asian: 100% ECOG PS=1: 37% Never smoker: 69% Stage IV: 48%	Gefitinib (n=86) Median OS: 34.8 m Median PFS: 9.2 m	Cisplatin+Docetaxel (n=86) Median OS: 37.3 m Median PFS: 6.3 m	D/C due to AEs: 16% SAEs: NR Tx-related deaths: 1
First-SIGNAL³² Median f/u: 34 m	Median age: 57; Asian: 100% ECOG PS=1: 68% Never smoker: 100% Stage IV: 90%	Gefitinib (n=26) ^β Median OS: 27.2 m Median PFS: 8.0 m	Cisplatin+Gemcitabine (n=16) ^{β} Median OS: 25.6 m Median PFS: 6.3 m	D/C due to AEs: NR AEs ≥ Grade 3: 29% Tx-related deaths: 0
EURTAC ^{44,45} Median f/u: 40.7 m (erlotinib) vs. 22.1 m (chemo)	Median age: 65; Asian: 0 ECOG PS=1: 53% Never smoker: 69% Stage IV: 92%	Erlotinib (n=86) Median OS: 22.9 m Median PFS: 10.4 m	Cisplatin+Gemcitabine/ Docetaxel (n=87) Median OS: 22.1 m Median PFS: 5.1 m	D/C due to AEs: 13% SAEs: 32% Tx-related deaths: 1
ENSURE ⁴⁶ Median f/u: 28.9 m (erlotinib) vs. 27.1 m (chemo)	Median age: 57; Asian: 100% ECOG PS=1: 79% Never smoker: 71% Stage IV: 92%	Erlotinib (n=110) Median OS: 26.3 m Median PFS: 11.0 m	Cisplatin+Gemcitabine (n=107) Median OS: 25.5 m Median PFS: 5.5 m	D/C due to AEs: 3% SAEs: 14% Tx-related deaths: NR
OPTIMAL^{47,48} Median f/u: 15.6 m	Median age: 58; Asian: 100% ECOG PS=0-1: 94% Never smoker: 71% Stage IV: 90%	Erlotinib (n=82) Median OS: 22.8 m Median PFS: 13.1 m	Carboplatin+Gemcitabine (n=72) Median OS: 27.2 m Median PFS: 4.6 m	D/C due to AEs: 1% SAEs: 12% Tx-related deaths: 0
TORCH³⁴ Median f/u: 24.3 m	Median age: 62; Asian: 3% ECOG PS=1: 50% Never smoker: 21% Stage IV: 89%	$\begin{array}{l} \mbox{Erlotinib} \\ (n=20)^{\beta} \\ \mbox{Median OS: 18.1 m} \\ \mbox{Median PFS: 9.7 m} \end{array}$	$\begin{array}{l} \textbf{Cisplatin+Gemcitabine} \\ (n=19)^{\beta} \\ \text{Median OS: 32.5 m} \\ \text{Median PFS: 6.9 m} \end{array}$	D/C due to AEs: NR SAEs: NR Tx-related deaths: NR

 α Outcomes from independent review; β EGFR+ subgroup, AEs reflect overall population; f/u= follow-up; NR=not reported; ECOG PS=Eastern Cooperative Oncology Group Performance Status; OS=overall survival; PFS=progression-free survival; HR=hazard ratio; ORR=objective response rate; D/C=discontinuation; AEs=adverse events; Tx=treatment; NR=not reported

Clinical Benefits

A detailed review of each clinical outcome of interest is presented in the sections that follow. All key studies were designed primarily to measure improvement in PFS, with the exception of the First-SIGNAL and TORCH trials of gefitinib and erlotinib, respectively, in which the primary endpoint was overall survival.

Overall Survival

RCTs had high rates of crossover and showed no OS benefit for TKIs compared with a platinumbased chemotherapy doublet. The most likely explanation for the lack of OS benefit seen in RCTs of TKIs with high rates of crossover is that treatment with a TKI improves survival whenever it is given in the sequence of therapy, such that patients randomized to chemotherapy who receive TKI therapy at the time of progression have similar survival in comparison to patients treated initially with a TKI. Observational data suggest that first-line TKI therapy as a class increases OS by approximately 8.9 months, although there is substantial uncertainty in this figure. Limitations in the evidence base preclude determining whether there are clinically important OS differences between the TKIs.

Improving overall survival (OS) and quality of life (QoL) are generally the patient-important goals of cancer therapy. Assessing the true survival benefit of an emerging therapy can be difficult when study participants are permitted to cross over to receive the alternative study treatment after tumor progression and the key studies included in the sample set for this review had high levels of crossovers (approximately 45 to 90% of patients in the chemotherapy arms crossed over to treatment with a TKI; see Appendix Table C3 for crossover rates). We present OS data here, noting that these results are likely biased estimates of the true survival benefit of the TKIs.

The ten randomized trials comparing a TKI with a platinum-based chemotherapy doublet all included data on OS, but no study showed statistical differences between the arms. Median OS was similar between TKIs, ranging from 18.1-26.3 months with erlotinib, 21.6-34.8 months with gefitinib, and 23.1-28.2 months with afatinib. Platinum doublet regimens produced comparable median survival results. Hazard ratios were also comparable across trials, ranging from 0.84-1.58 with confidence intervals that all crossed 1. These results are presented in Figure 5.

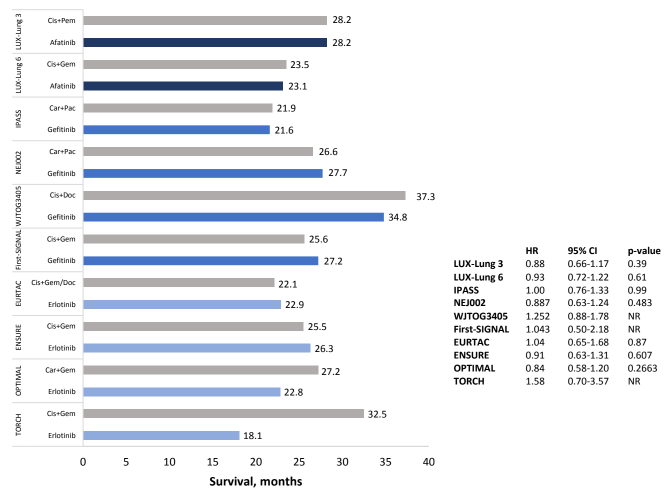


Figure 5. Overall survival: TKIs vs. platinum-based chemotherapy doublets

Cis=cisplatin; Car=carboplatin; pem=pemetrexed; gem=gemcitabine; pac=paclitaxel; doc=docetaxel

We identified only two studies evaluating differences between TKIs through direct comparison. One phase IIb randomized trial, LUX-Lung 7, compared afatinib and gefitinib and found no difference in OS (27.9 months vs. 25.0 months; hazard ratio [HR] 0.87, 95% CI 0.66-1.15; p=0.33).³⁸ The authors noted that survival data were not yet mature, with a median duration of follow-up of 27.3 months. A small observational study that compared gefitinib and erlotinib found no statistically significant difference in OS.⁴⁹

Given the paucity of head-to-head data comparing TKIs, we performed indirect comparisons of the TKIs using Bayesian network meta-analyses (NMAs). NMA was felt to be appropriate as the populations of the individual trials were sufficiently similar for comparisons of the relative effects of

TKI therapy compared with a platinum doublet across the trials. Detailed descriptions of methods and results can be found in Appendix D. Our results did not show statistical differences between agents, which aligns with the findings of previously published network meta-analyses (see Haspinger and Zhang in Appendix G for detailed descriptions of these studies).^{10,50}

We did not identify any subgroups for whom the TKIs had differential effects on overall survival. We also conducted meta-analyses stratified by the two common EGFR mutations (i.e., exon Del19 and L858R, respectively), and we did not find a differential effect of TKIs in these two subgroups (p=0.115, see Appendix E).

Challenges in Estimating Impact of TKIs on Overall Survival

As discussed above, the assessment of benefit of TKIs on OS is problematic due to high rates of crossover from chemotherapy to TKIs in the randomized trials, which is understandable for ethical reasons but has the potential to dilute estimates of treatment effect over longer periods of followup.^{35,51,52} PFS and OS appear to be substantially correlated in NSCLC in trials where patients rarely cross over, but are much less correlated in trials where crossing over is more common.⁵³ It appears most likely that the explanation for the lack of OS benefit seen in randomized trials of TKIs with high rates of crossover is that treatment with a TKI improves survival *whenever* it is given in the sequence of therapy, such that patients randomized to chemotherapy who receive TKI therapy at the time of progression have similar survival in comparison to patients treated initially with a TKI.

However, in estimating a potential incremental benefit of any of the TKI therapies in our analysis relative to treatment with a chemotherapy doublet alone, we need an estimate of the improvement in OS for the class. Several pieces of evidence inform such an estimate. For instance, historical survival rates in trials of platinum doublets are substantially lower than survival rates in both arms of trials of TKIs.^{35,51,52,54} Although suggestive, this may only reflect secular trends toward improved survival in patients with advanced NSCLC. In the OPTIMAL trial comparing erlotinib with carboplatin+gemcitabine, an analysis looked at OS in subgroups defined by post-trial therapy.⁵⁵ In the erlotinib arm, 30 patients received no post-trial therapy, and 1 patient received another TKI (TKI-only group); in the chemotherapy arm, 16 patients received no post-trial therapy and 4 patients received other chemotherapy (chemotherapy-only group). Median OS in the 31 patients who had only received TKI therapy was 20.7 months, and OS in the 20 patients who had received only chemotherapy was 11.2 months. It is likely, however, that these groups would not have had prognostic balance at baseline.

Within-trial comparisons offer a less biased evaluation of possible survival benefit with TKIs. A single RCT that compared gefitinib with carboplatin+paclitaxel, the IPASS trial, provided the best within-trial evidence that we found:⁵⁶

• There were 261 EGFR+ patients and 176 EGFR- patients

- Gefitinib treatment would not be expected to improve OS in patients who are EGFR-, as can be seen by its lack of benefit on PFS in EGFR- patients. In IPASS, for example, EGFR- patients had a significantly higher risk of progression when treated with gefitinib relative to chemotherapy (HR 2.85, 95% CI 2.05-3.98).⁵⁷
- As in other trials, treatment with gefitinib compared with a platinum doublet did not significantly improve OS among all patients (median OS 18.8 months vs. 17.4 months; HR 0.90, 95% CI 0.79-1.02), in the EGFR+ subgroup (21.6 months vs. 21.9 months; HR 1.00), or in the EGFR- subgroup (11.2 months vs. 12.7 months; HR 1.18).
- Comparing OS in EGFR+ patients in the TKI arm of the trial (21.6 months) with OS in EGFRpatients in the chemotherapy arm of the trial (12.7 months) can give us an estimate of the size of the improvement in OS with TKI therapy for EGFR+ patients, since EGFR- patients initially treated with chemotherapy would not be expected to benefit from crossing over to TKI therapy.
- For EGFR+ patients, this difference between arms (21.6 vs. 12.7 months) gives an *estimated median OS advantage with gefitinib compared with carboplatin/paclitaxel of 8.9 months*.
- Given the actual results seen in IPASS, this is a conservative estimate among the possible pairwise comparisons (as contrasted with, for instance, comparing the OS of all EGFR+ patients with OS of all EGFR- patients).

There are important reasons to be wary of this estimate of an 8.9 months survival advantage. It is controversial whether EGFR+ status is a marker for improved prognosis per se,⁵⁸⁻⁶⁰ and EGFR+ status may also be associated with improved survival because it is more common in non-smokers. Thus, the better survival seen in IPASS in EGFR+ patients could be due to EGFR status itself rather than the benefits of TKI therapy in EGFR+ patients. However, most estimates of any improved prognosis with EGFR+ tumors are small, and in IPASS, most patients were non-smokers, including 94% of EGFR+ patients and 90% of EGFR- patients.⁵⁶

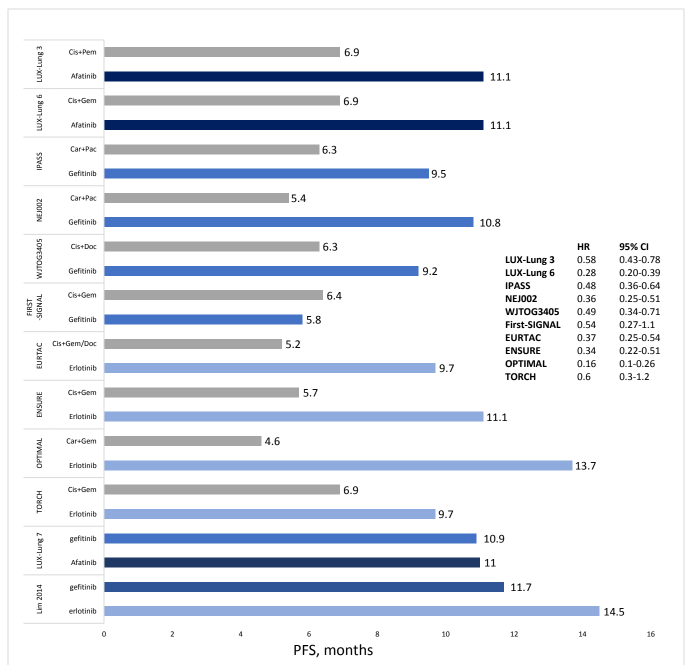
Although this estimate comes from data from a randomized trial, this is an observational post-hoc analysis of the results in subgroups of the original trial population. We are, however, unlikely to get direct trial evidence that provides the survival benefit of administering a TKI to patients with EGFR+ advanced NSCLC, as it would be felt to be unethical to withhold such therapy in a randomized trial.

Progression-free Survival

Evidence from RCTs indicates that all three TKIs provide statistically-significant improvements in PFS relative to platinum doublet chemotherapy, with HRs ranging from 0.30 to 0.45 (absolute improvements typically in the 3 to 6 month range). A head-to-head randomized trial, as well as

our network meta-analysis, provide evidence that treatment with afatinib likely provides a small benefit in PFS compared with gefitinib, however, PFS is mostly a surrogate endpoint.

All 11 key studies assessed PFS, ten of which compared a TKI to a platinum-based chemotherapy doublet, and one which compared afatinib to gefitinib. We identified an additional retrospective observational study⁴⁹ (described above) of gefitinib versus erlotinib. With the exception of the First-SIGNAL and TORCH trials of gefitinib and erlotinib, respectively, all key studies measured improvement in PFS as a primary endpoint. PFS is calculated from the time of the start of treatment to disease progression or death. Although it is a surrogate endpoint that is commonly considered in regulatory processes, the extent to which it correlates with overall survival varies, particularly when crossover rates are high (see discussion in "Challenges in Estimating Impact of TKIs on Overall Survival"). In general, TKIs improved PFS compared with platinum doublets, but there is insufficient evidence to distinguish the effects between individual TKIs. Figure 6 presents median PFS from these trials.





Cis=cisplatin; Car=carboplatin; pem=pemetrexed; gem=gemcitabine; pac=paclitaxel; doc=docetaxel

Only one phase IIb trial, LUX-Lung 7, directly compared two TKIs, afatinib and gefitinib. In this trial of 319 patients, those randomized to afatinib had a slightly longer median PFS but the treatment effect was statistically significant (11.0 months vs. 10.9 months; HR 0.73, 95% CI 0.57 to 0.95; p=0.017); although the difference in median PFS was only 0.1 months, the survival curves generally

showed greater separation after 12 months (Figure 7).³⁸ The previously-described retrospective observational study found no statistically significant difference in PFS with gefitinib or erlotinib.⁴⁹

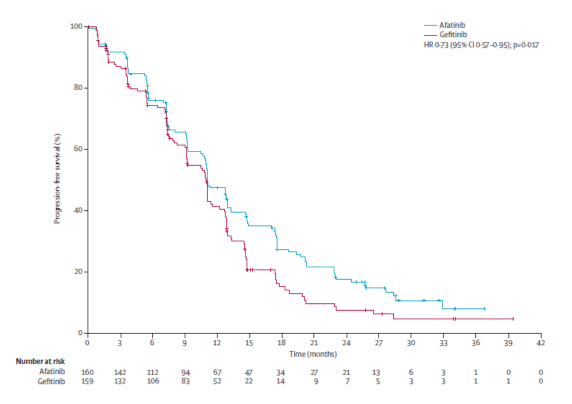


Figure 7. Progression-free survival curve in LUX-Lung 7

Source: Park K, Tan EH, O'Byrne K, et al. Afatinib versus gefitinib as first-line treatment of patients with EGFR mutation-positive non-small-cell lung cancer (LUX-Lung 7): A phase 2B, open-label, randomised controlled trial. The Lancet Oncology. 2016;17(5):577-589.

Compared with a platinum doublet, improvements in median PFS were similar across the trials of the TKIs (4-month benefit with afatinib, 3-5 months with gefitinib, and 3-9 months with erlotinib). A single trial of gefitinib versus cisplatin+gemcitabine (First-SIGNAL trial) did not find statistical differences in median PFS. The PFS curves crossed at seven months, with the gefitinib arm showing less progression during the first seven months but more progression afterwards.³² A single study (TORCH) of erlotinib versus cisplatin + gemcitabine failed to show statistical improvements in PFS in a subgroup analysis of 39 EGFR+ patients; this was likely due to the small sample size.³⁴ In the eight trials that found statistical improvements in PFS with TKIs, hazard ratios showed risk reductions for progression that ranged from 0.28-0.58 with afatinib, 0.30-0.49 with gefitinib, and 0.16-0.33 with erlotinib. Results from our network meta-analysis yielded hazard ratios of 0.38, 0.45, and 0.30 for afatinib, gefitinib, and erlotinib, respectively.

Our network meta-analysis combined both direct and indirect evidence on PFS for the TKIs, and showed no statistical differences among these agents. Results were similar when the network included all regimens of platinum-based doublets combined, or cisplatin-based and carboplatin-based regimens analyzed separately (Appendix D).

There is some evidence to suggest that the efficacy of TKIs varies slightly by type of EGFR mutation. A meta-analysis of seven trials that reported subgroup results for the two common types of EGFR mutations suggested that the type of mutation modified the effect of TKIs on PFS (p=0.004). Del 19 was associated with a greater PFS benefit than L858R (HR 0.28 [95% CrI 0.20-0.35] vs. 0.48 [0.36-0.43], see Appendix E).

Objective Response Rate

Evidence from randomized trials shows higher objective response rates (ORRs) with TKI therapy than with a platinum doublet. A head-to-head trial found a higher ORR with afatinib compared to gefitinib.

Figure 8 presents response data from each of the key trials. The majority of studies evaluated tumor response using RECIST criteria. Objective response was universally defined as a partial or complete response. Relative to therapy with a platinum doublet, a significantly greater proportion of patients had a partial response or better with the studied TKIs. Among the TKIs, the objective response rate (ORR) varied substantially, with no clear differences between agents: ORRs ranged from 56-67% with afatinib, 62-85% with gefitinib, and 42-83% with erlotinib. However, LUX-Lung 7 that compared afatinib and gefitinib head-to-head, found a significantly higher ORR with afatinib (70% vs. 56%; Odds ratio [OR] 1.87, 95% CI 1.18-2.99; p=0.0083).³⁸

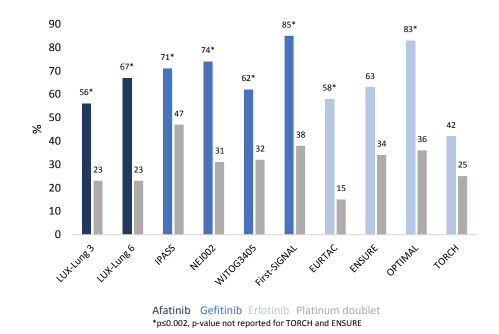


Figure 8. Objective response rate: TKIs vs. platinum-based chemotherapy doublets

Quality of Life

Quality of life (QoL) improvements were greater with TKI therapy than with a platinum doublet. Evidence is inadequate to distinguish QoL benefits among the TKIs.

We identified six RCTs, representing two trials for each individual TKI, that investigated the impact of TKIs on quality of life (QoL) in patients with NSCLC. The one head-to-head trial did not assess QoL. Comparisons across agents and trials were problematic, as different instruments were used for different drugs and clinically relevant improvement and worsening were defined accordingly. Overall, all six trials indicated that TKIs provided significant improvements relative to comparator treatment on at least one QoL outcome and five of them were clinically meaningful. Results are summarized in Table 4. Additional details are presented in Appendix C.

Trial name	QoL instrument	Domain	Improvement in mean score	Clinically meaningful improvement
LUX-Lung 3	QLQ C30	Global Health Status/QoL	Y	N
		Physical Functioning	Y	N
		Role Functioning	Υ	N
		Cognitive Functioning	Y	N
		Emotional Functioning	N	Ν
		Social Functioning	N	N
LUX-Lung 6	QLQ C30	Global Health Status/QoL	Υ	Y
		Physical Functioning	Y	Y
		Role Functioning	Y	Y
		Cognitive Functioning	Y	Υ
		Emotional Functioning	Y	Υ
		Social Functioning	Y	Υ
IPASS	FACT-L	Physical well-being Functional well-being Social well-being Emotional well-being LCS Relationship with doctor	Y	Y
	ТОІ	Physical well-being Functional well-being LCS	Y	Y
NEJ002	The Care Notebook	Physical well-being	Y	Ŷ
		Mental well-being	Ν	N
		Life well-being	Υ	Υ
ENSURE	FACT-L	See IPASS	NR	Y
	TOI	See IPASS	NR	Y
OPTIMAL	FACT-L	See IPASS	Y for subscales: Physical well-being Emotional well-being LCS	Y
	TOI	See IPASS	NR	Y

Table 4. Improvement with TKIs on QoL domains

Symptom Control

Symptom control was greater with TKI therapy than with a platinum doublet. Evidence is inadequate to distinguish symptom control benefits among the TKIs.

The same six trials that evaluated patient QoL also measured symptom control, using subscales from the QoL questionnaires. Again, all six studies showed that TKIs had a greater benefit on at least one symptom-related outcome. Symptoms that showed improvement with TKI therapy in at least one trial included dyspnea,⁶¹ pain, ⁶¹ and a composite symptom score that included shortness of breath, weight loss, clarity of thinking, cough, good appetite, chest tightness, and ease of

breathing.^{33,62} Symptoms that showed delayed deterioration with TKI therapy in at least one trial included dyspnea,^{61,63} cough,^{61,63} pain,⁶³ shortness of breath,⁶⁴ and a composite score.⁵²

Harms

Rash, diarrhea, and liver function abnormalities are the most common side effects with TKIs. All TKIs appear to be better tolerated than chemotherapy with platinum doublets, which have much higher rates of hematologic toxicity. Afatinib appears to have higher rates of diarrhea and rash, while liver function abnormalities are more common with gefitinib.

Adverse event (AE) frequencies and rates of Grade 3-4 (severe and life-threatening) events are reported by regimen in Table 5. The values in Table 5 represent weighted averages across trials. Relative to a platinum doublet, there were lower rates of discontinuation due to AEs and fewer AEs of grade 3 or greater with the TKIs. Platinum-based chemotherapy doublets were associated with a higher incidence of hematological AEs such as anemia and neutropenia, whereas the most common adverse events among the TKIs included dermatologic toxicity (skin rash), hepatotoxicity (elevated levels of aspartate aminotransferase and/or alanine aminotransferase), and diarrhea. Among TKIs, afatinib appears to have a higher incidence of diarrhea and rash, while a greater proportion of patients developed increased aminotransferase concentrations with gefitinib. These findings are consistent with previous systematic reviews, which are described in detail in Appendix G (Haspinger, Zhang).^{10,50}

%	Afatinibα	Gefitinib	Erlotinib	PBCD			
Treatment- related deaths	1	0	1	1			
D/C due to AEs	9	8	8	15			
≥Grade 3 AES	45	37	35	62			
	Grade 3-4 AEs, %						
Anemia	0	2	3	9			
Appetite loss	3	4	1	4			
Diarrhea	10	3	4	1			
Fatigue	2	6	7	9			
Increased ALT [‡]	3	18	2	2			
Increased AST	0	17	2	3			
Nausea	1	0	0	3			
Neutropenia	1	1	0	42			
Rash ^Ω	15	8	9	3			

Table 5. Grade 3-4 adverse events: TKIs

Values represent weighted averages across key trials; α 1 study only reported treatment-related AEs; \ddagger 3 studies reported elevated aminotransferase and were integrated into both ALT and AST calculations; Ω 3 studies included reported as acne/rash; D/C=discontinuation; AEs=adverse events; ALT=alanine aminotransferase; ASP=aspartate aminotransferase; PBCD=platinum-based chemotherapy doublets

Controversies and Uncertainties

There are relatively few head-to-head studies of TKIs, and our NMAs show wide credible intervals for the HR for PFS when comparing the three TKIs we evaluated for first-line therapy. As such, it is difficult to judge whether there are important clinical differences in the effectiveness of these therapies. The results of LUX-Lung 7 do suggest a PFS benefit of afatinib over gefitinib. In preliminary input, manufacturers questioned whether the unblinded nature of this trial and evaluation time bias (which can arise if the timing of assessment for progression has different patterns between the arms of the trial) might have influenced results, but progression was assessed in a blinded fashion and our NMAs show similar estimates of effect (trend toward PFS benefit for afatinib) regardless of whether LUX-Lung 7 was included in the network.

We also received preliminary input from manufacturers about the importance of considering the specific EGFR mutation when comparing agents across studies, but we found similar estimates of relative effect on PFS when we constructed networks for the two most common mutations individually. Other such input addressed the implications of the different populations being studied (predominantly Caucasian or Asian) on the results; however, we received input from various experts that once EGFR status is controlled for, ethnicity does not appear to be an effect modifier for TKI treatment.

The concerns mentioned above apply mainly to estimates of PFS, where benefits relative to platinum-based chemotherapy have been observed for each of the TKIs. Potentially more problematic issues for creating relative estimates of OS among the TKIs are the very wide credible intervals seen in the NMAs, and also the additional uncertainties generated by the effects of crossovers within the trials. As discussed previously, our largest uncertainties fall around the effect of TKIs as a class on OS, given that the randomized results within each of the trials were problematic due to the crossover effect. Had we used those results, we would have concluded that TKIs have no effect on OS. We received preliminary input from manufacturers suggesting that we use the OS benefit seen for afatinib in the exon 19 deletion subgroups of LUX-Lung 3 and LUX-Lung 6. If we used the point estimates from these trials, even in this subgroup we would have seen a smaller OS benefit than we chose to apply to the TKIs as a class, and would additionally have concluded that afatinib (and perhaps the other TKIs) substantially worsens survival in patients with the L858R mutation. This seems unlikely, given that a PFS benefit was observed in this subgroup. However, we do think the PFS results across all the TKI trials suggest that there may be a larger OS benefit with TKIs in patients with tumors with exon 19 deletions and a smaller OS benefit in tumors with the L858R mutation. We do not feel we have a good way to estimate this differential benefit, however, given the crossover concerns.

Although we are estimating the survival benefit conferred by adding TKI therapy to the prior standard of care of a platinum doublet, it is important to note that TKI treatment has already

become standard of care for patients with advanced EGFR+ NSCLC, and that those patients with the T790M mutation (about half of EGFR+ patients who progress) will also be treated at the time of progression with the TKI osimertinib.^{7,65} Second line TKI therapy in such patients likely provides additional survival benefits, although analysis of this type of treatment pathway was beyond the scope of our review. Similarly, although not commonly used as a comparator in the clinical trials, modern chemotherapy for adenocarcinoma would typically include maintenance treatment with pemetrexed, which has been shown to improve overall survival (OS).⁶⁶ However, given the evidence base, our analysis does not consider the incremental effectiveness nor the incremental cost of maintenance pemetrexed. (Separate from this use as maintenance therapy, pemetrexed is used as a component of many initial chemotherapy regimens, and we do consider this use in our analysis.)

Cisplatin appears to be slightly more effective than carboplatin, although it has not been a universal comparator in all relevant TKI trials.⁶⁷ Analyses that adjusted for this where possible gave similar results, and we received expert input that this difference in effectiveness is not felt to be an important concern in interpreting results of trials that used a chemotherapy doublet as a comparator.

Finally, we heard some concerns from patient groups that rash may have been underreported in the clinical trials of TKIs because of patient concerns that they might be taken off therapy if they accurately reported this adverse event. Although this is an important consideration in thinking about the estimates of adverse event rates, overall rates of all serious adverse events appear to be substantially lower with TKIs than with platinum doublets.

PD-1 Immunotherapies

Although our initial scope described three distinct populations of patients who might be treated with PD-1 immunotherapy (described above and below), all relevant evidence came from four randomized trials applicable to population P3. We discuss this population first in sequence, followed by discussions of populations P2 and P4, as well as our limited ability to extrapolate from population P3 to these other populations. Summaries of the results are provided in the sections discussing each population.

Our review of PD-1 immunotherapy in patients with advanced NSCLC focused on atezolizumab, nivolumab, and pembrolizumab in three distinct populations: in patients with a tumor that does not have a driver mutation and has progressed after first-line treatment with a platinum-based chemotherapy doublet (population P3); in patients with a tumor that does not have a driver mutation who have not previously been treated for advanced disease (population P2); in patients with an EGFR+ tumor that has progressed after first-line or first- and second-line TKI therapy (population P4). As described previously in Section 4.1, we assessed these therapies in relation to single-agent docetaxel in patients without a driver mutation, and relative to a platinum-based chemotherapy doublet in patients without a driver mutation who were not previously treated for advanced disease or with an EGFR+ tumor receiving second- or third-line treatment. The sections that follow discuss the overall survival, progression-free survival, response, quality of life, symptom control, and harms associated with these agents in each of the populations of interest.

Study Selection

Our literature search identified eight references that met our criteria for atezolizumab, nivolumab, or pembrolizumab; these citations related to one systematic review (see Appendix G) and four individual studies, all four of which were deemed good quality and compared a PD-1 immunotherapy to single-agent docetaxel. We identified one phase IIb RCT of atezolizumab, two phase III RCTs of nivolumab, and one phase II/III trial of pembrolizumab. In addition, three conference abstracts associated with the two nivolumab studies were included in our study set. All four trials provided evidence that informed our analysis of the use of PD-1 immunotherapy in EGFR-patients in the second-line setting (P3); we found no studies that focused on first-line use in EGFR-patients (P2) or on treatment of EGFR+ patients who had progressed after TKI therapy (P4). As discussed below, results from two trials of PD-1 immunotherapy in a first-line setting are expected to become available in the near future. Appendix Table C1 presents the number of studies identified for each population of focus.

Key Studies

We considered four RCTs to be key studies of interest for this review, which are summarized in Table 6. Important outcomes from each trial are also provided in Appendix Table B1, and described in further detail in the sections that follow.

Key Trials	Patient Characteristics	Treatment	Comparator	Harms (Treatment Arm)
CheckMate 017 (nivolumab)	Median age: 63 Asian: 2% ECOG PS=1: 76% Never smoker: 6% 1 prior therapy: 100% Non-squamous: 0 EGFR+: NR EGFR-: NR	Nivolumab (n=135) Min. follow-up: 11 m Overall Median OS: 9.2 m Median PFS: 3.5 m OS HR: 0.59 (95% CI 0.44-0.79) PFS HR: 0.62 (95% CI 0.47-0.81) EGFR+ (NR) EGFR- (NR)	Docetaxel (n=137) Median OS: 6.0 m Median PFS: 2.8 m	D/C due to TEAEs: 3% AE ≥ Grade 3: 7% Tx-related deaths: 0
CheckMate 057 (nivolumab)	Median age: 62 Asian: 3% ECOG PS=1: 69% Never smoker: 20% 1 prior therapy: 88% Non-squamous: 100% EGFR+: 14% EGFR-: 58%	Nivolumab (n=292) Docetaxel (n=290) Follow-up (OS): 13.2 m (Additional follow-up 17.2 m) Overall Median OS: 12.2 m Median OS: 39 m ^a 18-m OS: 23 m ^a Median PFS: 2.3 m Median PFS: 4.2 m OS HR: 0.72 (95% CI 0.60-0.88) PFS HR: 0.91 (95% CI 0.76-1.09) EGFR+ OS HR: 1.18 (95% CI 0.69-2.00) PFS HR: 1.46 (95% CI 0.90-2.37) EGFR- OS HR: 0.66 (95% CI 0.51-0.86) PFS HR: 0.83 (95% CI 0.65-1.06)		D/C due to TEAEs: 5% AE ≥ Grade 3: 46% Tx-related deaths: 1
KEYNOTE-010 (pembrolizumab)	Mean age: 63.0 Asian: 21% ECOG PS=1: 66% Never smoker: 19% 1 prior therapy: 70% Non-squamous: 70% EGFR+: 8% EGFR-: 85%	Pembrolizumab (n=344)∞ Docetaxel (n=343) Median follow-up: 13.1 m Docetaxel (n=343) Overall Median OS: 10.4 m Median PFS: 3.9 m Median PFS: 4.0 m OS HR: 0.71 (95% CI 0.58-0.88) PFS HR: 0.88 (0.74-1.05) EGFR+ OS HR: 0.88 (95% CI 0.45-1.70) PFS HR: 1.79 (95% CI 0.94-3.42) EGFR- OS HR: 0.66 (95% CI 0.55-0.80) PFS HR: 0.83 (95% CI 0.71-0.98)		D/C due to AEs: 10% AE ≥ Grade 3: 13% Tx-related deaths: 3
POPLAR (atezolizumab)	Median age: 62 Asian: NR ECOG PS=1: 68% Never smoker: 20% 1 prior therapy: 66% Non-squamous: 66% EGFR+: 11% ⁿ EGFR-: 89% ⁿ	Atezolizumab (n=144) Median follow-up: 14.8 m Overall Median OS: 12.6 m Median PFS: 2.7 m OS HR: 0.73 (95% Cl 0.53-0.99) PFS HR: 0.94 (95% Cl 0.72-1.23) EGFR+ (NR) EGFR- (NR)	Docetaxel (n=143) Median follow-up: 15.7 m Median OS: 9.7 m Median PFS: 3.0 m	D/C due to AEs: 8% AE ≥ Grade 3: 40% Tx-related deaths: NR

Table 6. Key trials of PD-1 immunotherapies

 Ω Of 83 patients with known EGFR status; α Overall survival with extended follow-up at 18 months; ∞ KEYNOTE-010 included two dosing groups, however we only report results from the 2mg/kg group as this dose is consistent with pembrolizumab's FDA prescribing information at the time of publication; NR=not reported; ECOG PS=Eastern Cooperative Oncology Group Performance Status; OS=overall survival; PFS=progression-free survival; HR=hazard ratio; D/C=discontinuation; AEs=adverse events; Tx=treatment

All four studies corresponded to our third population of interest (i.e., patients without a driver mutation who had disease recurrence or progression after treatment with a platinum-based doublet chemotherapy regimen). These studies were not exclusively composed of patients with a tumor without a driver mutation; however, our intended focus was to split out patients with and without an EGFR+ tumor because of concerns of effect modification by EGFR status. Even in this subgroup (patients with an EGFR- tumor), only two of the key studies (KEYNOTE-010 trial of pembrolizumab; CheckMate 057 trial of nivolumab) presented subgroup analyses in these patients.^{17,68} We present these results; we also present results from the overall populations of the remaining two studies, as they likely approximate the efficacy of the therapies in patients with an EGFR- tumor. One of the remaining studies (Checkmate 017 trial of nivolumab) was composed entirely of patients with squamous-cell NSCLC, who were not likely to have an EGFR+ tumor,⁶⁸ while 89% of patients in the final key study (POPLAR trial of atezolizumab) had an EGFR- tumor.²¹ Although the KEYNOTE-010 trial assessed relevant outcomes at two different doses of pembrolizumab (2 mg/kg and 10 mg/kg), we report results only from the 2 mg/kg group, as this dose is consistent with FDA's labeled indication for pembrolizumab at the time of this report's publication; subgroup analyses used combined results.

The trials specified similar inclusion criteria and trial populations were similar with respect to age and ECOG performance status. The proportion of never smokers was around 20% in all trials except for the CheckMate 017 trial of nivolumab (6%), where all patients had squamous NSCLC, and were thus more likely to have a history of smoking. The percentages of EGFR+ patients were generally low in these trials, ranging from 8% to 14% when reported. EGFR mutations are uncommon in squamous NSCLC, and the frequency was not reported in CheckMate 017. According to our scope, we focused on reporting results in EGFR- patients whenever possible and describe results in the population as a whole as an approximation when necessary, given the small proportion of EGFR+ patients in the trials.

All patients had provided tumor specimens for PD-L1 testing. However, PD-L1 expression levels were not comparable among trials because the investigators used different testing methods and cut-offs. For this same reason, subgroup analyses stratified by PD-L1 expression level were not comparable across drugs even at the same cut point. Furthermore, the KEYNOTE-010 trial was restricted to patients with at least 1% PD-L1 expression in tumor cells and provided no data on the effectiveness of pembrolizumab in PD-L1-negative patients.

Clinical Benefits

A detailed review of each clinical outcome of interest is presented in the sections that follow. All key studies were designed to measure improvement in overall survival as the primary outcome.

PD-1 Immunotherapies for Patients without a Driver Mutation who have Progressed after Treatment with a Platinum Doublet

<u>Summary</u>

- Trials of PD-1 immunotherapies used different assays to measure PD-L1 levels and had different PD-L1 cut-points both as entry criteria and for subgroup analyses. Given the difficulties in comparing results across trials and patient populations, we found inadequate evidence to distinguish among the PD-1 immunotherapies on any outcome.
- Evidence from RCTs indicates that in patients with advanced NSCLC without a driver mutation who have progressed after treatment with a platinum doublet, PD-1 immunotherapies improve survival compared with docetaxel.
- Patients with tumors that express high levels of PD-L1 are more likely to respond to PD-1 immunotherapies. However, only a minority of patients overall respond to these agents, even among those with high PD-L1 levels on assays. Conversely, even with negative PD-L1 level results, some patients do respond to PD-1 immunotherapy.
- Because of the limited follow-up in the existing studies, we are uncertain of how large the benefit is for the minority of patients who do respond to these agents.
- The most common adverse events seen with PD-1 immunotherapies are fatigue, nausea, and decreased appetite. Serious immune-related adverse events, including pneumonitis and encephalitis, can occur with these agents; these adverse events are not typically seen with chemotherapy. Overall, however, PD-1 immunotherapy is better tolerated than docetaxel.
- Evidence was inadequate to evaluate improvements in QoL with PD-1 immunotherapies.

Even with uncertainties about the duration of benefit with PD-1 immunotherapies, the current evidence base gives us high certainty that a substantial minority of patients with EGFR- advanced NSCLC do respond and achieve important gains in overall survival ("A").

Overall Survival

PD-1 immunotherapies improve survival overall compared with docetaxel. This improvement reflects prolonged benefits in a minority of patients and no benefit in the majority of patients, making standard descriptive statistics of survival benefit (median survival and hazard ratios) potentially misleading in understanding the overall effects of these therapies. Additionally, trials were not long enough to fully assess the survival benefit in patients who do respond to PD-1 immunotherapy. Higher levels of PD-L1 are associated with higher levels of response to PD-1

immunotherapies. The lack of comparability of the study populations precludes determining whether there are clinically important OS differences between the PD-1 immunotherapies.

All four key studies of PD-1 immunotherapies evaluated overall survival, although only two studies stratified results by mutation status. Median OS was not reported in the EGFR- subgroup, however statistically significant risk reductions indicate that the PD-1 immunotherapies provided a survival benefit over docetaxel in patients without an EGFR+ tumor. In the KEYNOTE-010 trial, pembrolizumab (2mg/kg and 10mg/kg groups combined) showed improved survival relative to single-agent docetaxel (HR 0.66, 95% CI 0.55-0.80). EGFR- patients treated with nivolumab in the CheckMate 057 trial saw a similar benefit (HR 0.66, 95% CI 0.51-0.86). Although the key trials of atezolizumab and nivolumab in squamous-cell carcinoma only reported OS in the overall populations, risk reductions were similar to those seen in EGFR- patients in the KEYNOTE-010 and CheckMate 057 trials: overall survival favored both atezolizumab (HR 0.73, 95% CI 0.53-0.99; p=0.040) and nivolumab in squamous-cell carcinoma (HR 0.59, 95% CI 0.44-0.78) relative to single-agent docetaxel. Absolute improvements in median OS were 2-3 months during the time of primary analysis. Extended follow-up in the CheckMate 057 showed a 16-month gain in overall survival with nivolumab for patients with non-squamous histology. These results are presented in Table 7.

	CheckMate 017		CheckMate 057		KEYNOTE-010		POPLAR	
	NIVO	DOCX	NIVO	DOCX	PEMB	DOCX	ATEZ	DOCX
Overall population								
Median OS, m (95% Cl)	9.2 (7.3-13.3)	6.0 (5.1-7.3)	12.2 (9.7-15.0) Extended f/u: 39 (34-45)	9.4 (8.1-10.7) Extended f/u: 23 (19-28)	10.4 (9.4-11.9)	8.5 (7.5-9.8)	12.6 (9.7-16.4)	9.7 (8.6-12.0)
HR (95% CI)	0.59 (0.4	14-0.79)	0.72 (0.6	0-0.88)	0.71 (0.58-0.88)		0.73 (0.53-0.99)	
EGFR- population								
HR (95% CI)	N	R	0.66 (0.5	1-0.86)	0.66 (0.55-0.80)		NR	

Table 7. Overall survival: PD-1 immunotherapy vs. docetaxel

NIVO=nivolumab; DOCX=docetaxel; PEMB=pembrolizumab; ATEZ=atezolizumab; OS=overall survival; m=months; HR=hazard ratio; f/u=follow-up; NR=not reported

Despite statistical risk reductions and clinically-significant absolute survival gains, we have substantial uncertainty about the true survival benefits of PD-1 immunotherapy over time. It is probable that the survival curves do not show proportional hazards, the assumption that underlies one of the most common ways of modeling and reporting relative benefits in survival analysis. This is demonstrated by OS curves that cross at 6 months in the CheckMate 057 trial of nivolumab and a general observation that OS curves flattened out after 15-18 months in the PD-1 immunotherapy arms in all of the key trials. To address this concern, we conducted a network meta-analysis of

parametric survival curves using Bayesian methods to capture time-varying HRs for OS. The analysis showed a trend toward decreasing median HRs over time for two of the agents (nivolumab: from 0.81 at 3 months to 0.46 at 18 months; atezolizumab: from 0.85 at 3 months to 0.51 at 18 months). The median HR with pembrolizumab stayed constant at 0.64 to 0.65 over time. Due to the small number of trials in the network, and the small number of events at each time point, all the credible intervals around the time-varying HR point estimates crossed 1.0 and overlapped with each other. Therefore, while evidence indicates that all of the PD-1 immunotherapies provide a significant OS benefit versus docetaxel, we have uncertainty about the true magnitude of this effect (Appendix D).

As discussed in the Controversies and Uncertainties section on page 50, notwithstanding our concerns about the proportional hazards assumption, it appears that there is a subpopulation of patients with NSCLC who have clinically important, durable responses to PD-1 immunotherapy. We explored PD-L1 expression and histology as two possible predictors of efficacy in patients treated with PD-1 immunotherapy. Also as discussed in that section, studies employed different thresholds and assays for measuring PD-L1 expression, making it difficult to draw conclusions across agents. However, subgroup analyses of three studies suggest that higher levels of PD-L1 expression correlate with better overall survival; the fourth study, the CheckMate 017 trial of nivolumab in patients with squamous NSCLC, did not show a consistent association between PD-L1 expression and survival. Subgroup analyses by PD-L1 expression level (not specific to EGFR- patients) are presented in Table 8.

	PD-L1 Expression Threshold	HR (95% CI)	PD-L1 Expression Threshold	HR (95% CI)
Nivolumab	<10%	0.70 (0.48-1.01)	≥10%	0.50 (0.28-0.89)
(CheckMate 017)	<1%	0.58 (0.37-0.92)	≥1%	0.69 (0.45-1.05)
Nivolumab	<10%	1.00 (0.76-1.31)	≥10%	0.40 (0.26-0.59)
(CheckMate 057)	<1%	0.90 (0.66-1.24)	≥1%	0.59 (0.43-0.82)
Pembrolizumab (KEYNOTE-010)	1-49%	0.76 (0.60-0.96)	≥50%	0.53 (0.40-0.70)
Atezolizumab	<median expression<="" th=""><th>1.1 (0.63-1.93)</th><th>≥50%</th><th>0.49 (0.22-1.07)</th></median>	1.1 (0.63-1.93)	≥50%	0.49 (0.22-1.07)
(POPLAR)	<1%	1.04 (0.62-1.75)	≥1%	0.59 (0.40-0.85)

We also examined whether PD-1 immunotherapies have differential efficacy according to histological diagnosis. In the POPLAR and KEYNOTE-010 trials of atezolizumab and pembrolizumab, respectively, hazard ratios for OS with PD-1 immunotherapy compared with docetaxel were somewhat *lower* in patients with non-squamous NSCLC (atezolizumab HR 0.69 [95% CI 0.47-1.01]; pembrolizumab HR 0.63 [95% CI 0.50-0.79]) than in patients with squamous NSCLC (atezolizumab

HR 0.80 [95% CI 0.49-1.30]; pembrolizumab HR 0.74 [95% CI 0.50-1.09]). In contrast, in the two CheckMate trials, which evaluated nivolumab in squamous- and non-squamous cell histologies, a *higher* hazard ratio was seen in the study of patients with non-squamous NSCLC (HR 0.72 [95% CI HR 0.60-0.88] vs. HR 0.59 [95% CI 0.44-0.79]). These results are reported in Table C4 of Appendix C. Subgroup meta-analyses of these data suggested that histology (squamous vs. non-squamous) was not an effect modifier for OS (p=0.847, see Appendix E).

Even though the use of PD-1 immunotherapy in first-line EGFR+ NSCLC was not in our scope, we compared EGFR+ and EGFR- groups in the meta-analysis, expecting it to inform our evaluation of second- or third-line use of immunotherapies in EGFR+ NSCLC. The results showed an OS benefit in the EGFR- subgroup (HR 0.66, 95% CI 0.58-0.74) but no benefit, in the EGFR+ subgroup (HR 1.12, 95% CI 0.69-1.81); the test for interaction with EGFR status was statistically-significant (p=0.036).

Progression-free Survival

PD-1 immunotherapies show mixed results on PFS compared with docetaxel. As with OS, this may represent effects in a mixed population of responders and non-responders; patients with tumors that express high levels of PD-L1 are more likely to have improvements in PFS with PD-1 immunotherapies.

All four key studies also evaluated PFS and found mixed results with regard to benefit, as well as the predictive value of PD-L1 expression level on the magnitude of benefit. The results are presented in Table 9.

The CheckMate 017 and CheckMate 057 trials compared nivolumab to docetaxel in patients with squamous and non-squamous NSCLC, respectively. In CheckMate 017, nivolumab improved median PFS (3.5 months vs. 2.8 months (HR 0.62, 95% CI 0.47 to 0.81; p<0.001).⁶⁹ Among the 225 (83%) patients with quantifiable PD-L1 expression at baseline, PD-L1 expression level was not found to be predictive of PFS. In contrast, CheckMate 057 found no difference in PFS overall; however, among 455 (78%) patients with quantifiable PD-L1 expression, higher PD-L1 expression was associated with an improvement in PFS with nivolumab at all three pre-specified levels (HR 0.70 for \geq 1% vs. 0.54 for \geq 5% vs. 0.52 for \geq 10%).⁶⁸

In KEYNOTE-010, EGFR- patients treated with pembrolizumab (2mg/kg and 10mg/kg groups combined) had a small PFS benefit (HR 0.83, 95% CI 0.71 to 0.98). In the total study population (EGFR+ and EGFR- patients), PFS was improved with pembrolizumab 2 mg/kg compared with docetaxel in patients with tumors with PD-L1 expression \geq 50% (median PFS 5.0 months vs. 4.1 months; HR 0.59, 95% CI 0.44 to 0.78; p=0.0001). PFS did not differ by treatment in patients with lower PD-L1 expression.

In POPLAR, there was no statistically significant difference in median PFS between arms, either in the total population²¹ or subgroups with different PD-L1 expression levels. There was, however, a non-significant trend toward greater PFS benefit with higher PD-L1 expression level (see Table 9).

	PD-L1 Expression	HR (95% CI)	Median PFS (mo	onths)
	Threshold	HK (95% CI)	PD-1 immunotherapy	Docetaxel
	≥10%	0.58 (0.33-1.0)	3.7	3.3
Nivolumab	≥5%	0.54 (0.32-0.90)	4.8	3.1
(CheckMate 017)	≥1%	0.67 (0.44-1.0)	3.3	2.8
	<1%	0.66 (0.43-1.0)	3.1	3.0
	≥10%	0.52 (0.37-0.75)	5.0	3.7
Nivolumab	≥5%	0.54 (0.39-0.76)	5.0	3.8
(CheckMate 057)	≥1%	0.70 (0.53-0.94)	4.2	4.5
	<1%	1.19 (0.88-1.61)	2.1	3.6
Pembrolizumab	≥50%	0.59 (0.45-0.78)	5.0	4.1
(KEYNOTE-010)	≥1%	0.88 (0.74-1.05)	3.9	4.0
	TC3 or IC3	0.60 (0.31-1.16)	7.8	3.9
Atezolizumab	TC2/3 or IC2/3	0.72 (0.47-1.10)	3.4	2.8
(POPLAR)*	TC1/2/3 or IC1/2/3	0.85 (0.63-1.16)	2.8	3.0
	TC0 and IC0	1.12 (0.72-1.77)	1.7	4.1

Table 9. Progression-free survival according to PD-L1 expression level (vs. docetaxel)

*TCO-3: percentages of tumor cells expression PD-L1 <1% (TCO), \geq 1% and <5% (TC1), \geq 5% and <50% (TC2), and \geq 50% (TC3); ICO-3: percentages of tumor area occupied by PD-L1-positive tumor infiltrating immune cells <1% (ICO), \geq 1% and <5% (IC1), \geq 5% and <10% (IC2), and \geq 10% (IC3).

As with OS, we conducted a network meta-analysis using parametric survival curves for PFS due to the concern that the proportional hazards assumption was violated. Our meta-analysis of nivolumab and atezolizumab showed no PFS benefit at one month but hazard ratios that gradually improved over time; a similar trend was seen with pembrolizumab. Due to the small number of trials, almost all credible intervals crossed one and heavily overlapped. (Appendix D).

We also used HRs to explore subgroup effects when survival curves were not available. Subgroup meta-analysis using HRs suggested that histology (squamous vs. non-squamous NSCLC) was not an effect modifier for PFS (p=0.104, see Appendix E). Our meta-analysis of EGFR- and EGFR+ subgroups suggested that EGFR status *is* an effect modifier for PFS (p=0.002) with a PFS benefit in EGFR- patients (HR 0.80, 95% CI 0.72-0.90) and an *increased* risk of progression or death in EGFR+ patients (HR 1.57, 95% CI 1.07-2.31).

Objective Response Rate

Objective response rates (ORRs) and duration of response provide another way to assess the benefits of PD-1 immunotherapies, given the difficulties with interpreting median survival and hazard ratios for OS. Patients with tumors that expressed high levels of PD-L1 had significantly

higher ORRs with PD-1 immunotherapies than docetaxel, and duration of response was much greater with PD-1 immunotherapies. In some trials, among patients who had responded to PD-1 immunotherapy, the duration of response at the time of trial completion could not be reported as a median number because the halfway point for that duration had not been reached.

Treatment response was evaluated in each of the key PD-1 immunotherapy studies, although no study stratified response endpoints by EGFR mutation status. Nevertheless, the majority of patients participating in the trials of focus did not have a driver mutation, so results from the ITT population may indicate what the response would look like in patients without an EGFR+ tumor.

Table 10 presents response data from the key studies. Objective response rate (ORR) was universally defined as a partial response or better using RECIST v1.1 criteria. Nivolumab- and pembrolizumab-treated patients had significantly higher rates of response relative to docetaxel (18-20% vs. 9-12%). Moreover, the responses were durable and still ongoing at the time of analysis. While atezolizumab and docetaxel produced almost identical ORRs (15%), the median duration of response was about seven months longer with atezolizumab.

Interestingly, while nivolumab and pembrolizumab were associated with higher rates of objective response relative to docetaxel, patients treated with these agents also had more progressive disease. This phenomenon may indicate that PD-1 immunotherapies are best suited for patients with certain clinical characteristics, such as a higher level of PD-L1 expression (see Table 11). As with overall survival, objective response rates among those treated with PD-1 immunotherapy were higher in the subgroups that had greater levels of PD-L1 expression. The CheckMate 017 trial, which evaluated nivolumab in patients with advanced squamous-cell NSCLC, was the only key trial that did not show an association between PD-L1 expression and response.

	CheckMa	te 017 ⁷⁰	CheckMa	ate 057 ⁶⁸	KEYNOT	E-010 ¹⁷	POP	LAR ²¹
	NIVO	DOCX	NIVO	DOCX	PEMB	DOCX	ATEZ	DOCX
Objective Response, % (95% Cl)	20.0 (14-28)	8.8 (5-15)	19.2 (15-24)	12.4 (9-17)	18.0 (14.1-22.5)	9.3 (6.5-12.9)	14.6 (NR)	14.7 (NR)
Odds ratio (95% Cl)	2.6 (1.3-5.5)		1.7 (1.1-2.6)		NR		NR	
p-value	0.008		0.02		0.0005		NR	
Median time to response, months (range)	2.2 (1.6-11.8)	2.1 (1.8-9.5)	2.1 (1.2-8.6)	2.6 (1.4-6.3)	2.1 (2.1-4.1)	2.1 (2.1-4.1)	NR	NR
Median duration of response, months (range)	Not reached (2.9-20.5+)	8.4 (1.4-15.2+)	17.2 (1.8-22.6+)	5.6 (1.2-15.2+)	Not reached (4.2-12.5)	6 (2.7-6.1)	14.3 (11.6-NE)	7.2 (5.6-12.5)
Progressive disease, n (%)	56 (41)	48 (35)	129 (44)	85 (29)	124 (37) ^α	89 (29) ^α	NR	NR

+ indicates a censored value due to ongoing response at time of analysis; α discontinued treatment due to progressive disease; ATEZ=atezolizumab; DOCX=docetaxel; NIVO=nivolumab; PEMB=pembrolizumab; NR=not reported; NE=non-estimable

	PD-L1 Expression Threshold	PD-1 immunotherapy ORR (%)	Docetaxel ORR (%)
Nivolumab	≥10%	19	9
(CheckMate 017)	<1%	17	10
	Total population	20.0	8.8
Although succession	≥10%	37	13
Nivolumab (CheckMate 057)	<1%	9	15
(Checkiviate 057)	Total population	19.2	12.4
Pembrolizumab	≥50%	30.2	7.9
(KEYNOTE-010)	Total population	18.0	9.3
	≥50%	37.5	13.0
Atezolizumab (POPLAR)	<1%	7.8	9.8
	Total population	14.6	14.7

Table 11. Objective response according to PD-L1 expression level (vs. docetaxel)

Quality of Life

Evidence was inadequate to assess the effects of PD-1 immunotherapies compared with docetaxel on quality of life.

Only one of the four trials evaluated QoL, and this showed improvement with nivolumab, although the results have only appeared in an abstract. CheckMate 017 assessed patient-reported health status using the EQ-5D preference-based health state utility measure (EQ-5D index; scaled from 0-1) and visual analog scale (EQ-VAS; scaled from 0-100). The minimum clinically-important difference (MID) is defined as 0.08 for the EQ-5D index and 7 for the EQ-VAS.⁷¹ While both EQ-5D and EQ-VAS were statistically significantly higher during a 48-week and 54-week follow-up than at baseline in the nivolumab arm ($p\leq0.05$), they were not different from baseline in the docetaxel arm at week 18, after which the sample size dropped below 10 and no analysis was conducted. EQ-VAS was found to have a statistically and clinically significant deterioration at the first follow-up after treatment discontinuation in the docetaxel arm, while no deterioration was observed in the nivolumab arm. Statistical tests comparing the nivolumab and docetaxel arms were not reported.

Symptom Control

Limited evidence showed no benefits of PD-1 immunotherapies on symptom control compared with docetaxel.

Two trials assessed symptom improvement with nivolumab. The evidence was insufficient to show any benefit with nivolumab on symptom control. Both CheckMate 017 and CheckMate 057 trials measured symptom burden using the LCSS average symptom burden index (LCSS ASBI), computed

by averaging 6 individual symptom scores (anorexia, fatigue, cough, dyspnea, hemoptysis, and pain). The minimally important difference (MID) was defined as a change of \geq 10 points on a 0-100 scale. At 27-week follow-up in CheckMate 057, symptom improvement rates were similar in the nivolumab and docetaxel arms (17.8% vs. 19.7%), and no changes in mean LCSS ASBI scores exceeded the MID in either arm.⁷² Similarly, CheckMate 017 found little difference in symptom improvement at week 12 (20.0% vs. 21.9%).⁷³

Harms

Fatigue, nausea, and loss of appetite are the most common adverse events seen with PD-1 immunotherapies. Immune-related adverse events can occur and can affect various organs, including the lungs, brain, liver, and skin. Some of these events can be severe. Overall, severe adverse events, primarily hematologic adverse events, are more common with docetaxel than with PD-1 immunotherapy.

Treatment emergent adverse event (TEAE) frequencies and rates of Grade 3-4 (severe and lifethreatening) events are reported by regimen in Table 12. We did not identify a single study that stratified safety outcomes by EGFR mutation status and therefore report data from the overall study populations.

The PD-1 immunotherapies were well tolerated in the key trials, with safety profiles that were generally superior to docetaxel. Patients treated with atezolizumab, nivolumab, or pembrolizumab had lower rates of discontinuation due to TEAEs, fewer grade 3-4 TEAEs, and fewer treatment-related deaths compared to patients treated with docetaxel. The higher rates of grade 3-4 TEAEs observed with docetaxel were attributable mainly to hematologic toxicity.

	POPLARα		CheckN	late 017	Check	Mate 057	KEYNOTE-010 ^β	
	ATEZ	DOCX	NIVO	DOCX	NIVO	DOCX	PEMB	DOCX
Treatment-related deaths, n (%)	1 (0.7)	3 (2)	0	3 (2)	1 (0.3)	1 (0.3)	3 (0.9)	5 (2)
D/C due to TEAEs %	1	18	3	10	5	15	4	10
≥Grade 3 TEAES, %	11	39	7	55	10	54	13	35
				Grade 3-	4 AEs, %			
Alopecia	0	1	0	1	0	0	0	1
Anemia	NR	NR	0	3	<1	3	1	2
Asthenia	1	3	0	4	<1	2	<1	2
Decreased appetite	2	0	1	1	0	1	1	1
Diarrhea	1	4	0	2	1	1	1	2
Dyspnea	7	2	NR	NR	<1	0	0.6	1.3
Fatigue	NR	NR	1	8	1	5	1	4
Hypothyroidism	1	0	0	0	0	0	0	0
Musculoskeletal pain	2	2	0	0	NR	NR	0	0
Myalgia	1	3	0	0	<1	0	0	0
Nausea	1	0	0	2	1	1	<1	<1
Leukopenia	NR	NR	1	4	0	8	0	2.6
Neutropenia	0	12	0	30	<1	27	0	12
Febrile Neutropenia	0	8	0	10	0	10	0	4.9
Peripheral neuropathy	0	1	0	2	0	1	0	0.3
Pneumonia	6	2	0	0	0	2	0.9	1.3
Pneumonitis	NR	NR	0	0	1	<1	1.8	0.3
Rash	NR	NR	0	2	<1	0	<1	0

Table 12. Grade 3-4 treatment-related harms: PD-1 immunotherapy

 α POPLAR AEs are all-cause. Values were estimated from chart; β Grade 3-5 TEAEs, PEMB results are reported from treatment arm that received 2 mg/kg; AE=Adverse event; D/C=discontinuation; TEAEs=treatment emergent adverse events; ATEZ=atezolizumab; DOCX=docetaxel; NIVO=nivolumab; PEMB=pembrolizumab; NR=not reported

PD-1 immunotherapies have been associated with immune-related adverse events (irAEs), which may include dermatologic toxicity (e.g., rash, pruritus), diarrhea or colitis, hepatotoxicity (elevations in serum levels of aspartate aminotransferase [AST] and alanine aminotransferase [ALT]), pulmonary inflammatory complications (e.g., pneumonitis, pneumonia), and endocrinopathies (e.g., hypothyroidism).⁷⁴ The most common adverse events seen with PD-1 immunotherapies are fatigue, nausea, and decreased appetite. Table 13 presents the most frequently-reported treatment-related AEs of any grade of severity, as well as AEs with possible immune etiology. Immune-mediated TEAEs occurred with greater frequency with the PD-1 immunotherapies relative to docetaxel.

	POP	LAR ^α	CheckN	late 017	CheckN	late 057	KEYNO	TE-010
%	ATEZ	DOCX	NIVO	DOCX	NIVO	DOCX	PEMB	DOCX
Fatigue	20	35	16	33	16	29	14	25
Nausea	12	27	9	23	12	26	11	15
Decreased appetite	18	16	11	19	10	16	14	16
Asthenia	6	13	10	14	10	18	6	11
Diarrhea	7	22	8	20	8	23	7	18
	Immune-mediated TEAS							
Pruritus	NR	NR	2	0	8	1	7	2
Rash	NR	NR	4	6	9	3	9	5
Hypothyroidism	6	0	4	0	7	0	7	<1
Pneumonitis	3	NR	5	0	3	1	5	2
AST	4	NR	2	1	3	1	3	1
ALT	4	NR	2	1	3	1	5	1
Colitis	NR	NR	1	0	1	0	1	0

Table 13. Common treatment-related adverse events with PD-1 immunotherapy (any grade) andTEAEs of immune etiology

 α POPLAR values were estimated from chart; hypothyroidism and immune-mediated TEAEs not specifically treatment-related; TEAE=treatment emergent adverse events; AST=aspartate aminotransferase; ALT=alanine aminotransferase; NR=not reported

Controversies and Uncertainties

Although the comparator in all four trials of PD-1 immunotherapies was docetaxel, we found no head-to-head trials comparing these agents. As such, it is difficult to assess whether there are any important differences in outcomes with the three agents we evaluated. Two of the agents, nivolumab and pembrolizumab are directed at PD-1; atezolizumab is directed at PD-L1. Although this different target might argue for considering atezolizumab separately, our meta-analysis suggests little heterogeneity of effect across these three agents.

There are some concerns about appropriate assessment of progression with immunotherapies. There may be a phenomenon of "pseudoprogression", where the enhanced immune reaction with therapy can lead to lesions appearing to have progressed on imaging even if no actual progression has occurred. Although this has been felt to be important in assessing response to immunotherapies with other tumor types, clinical experts had differing opinions as to whether this is an important issue in NSCLC.

We received a number of comments regarding how to appropriately summarize the effects of PD-1 immunotherapy, as survival curves suggest that the proportional hazards assumption may not be valid, and that there may be a long survival tail among responders to therapy. Our analysis did suggest violations of proportional hazards, particularly for the trials of atezolizumab and nivolumab. It seems likely that the difficulty in using a proportional hazards model is generated by two populations in the PD-1 immunotherapy arms of the trials: a majority of patients who do not have sustained responses to therapy and have a high hazard for progression/mortality, and a minority of patients who do have sustained responses and have a much lower hazard. This is also reflected in

the median duration of response results seen in the trials (Table 9). There are relatively few data to allow assessment of whether there is a very long tail of responders beyond two years, but this is clearly an important issue in understanding the potential benefit of PD-1 immunotherapy.

We received comments that squamous and non-squamous histologies should be analyzed separately, given the apparently different responses to nivolumab in CheckMate 017 (squamous histology) and CheckMate 057 (non-squamous histology). However, as discussed above, we found no consistent direction of differences across the three PD-1 immunotherapies when we looked at the squamous and non-squamous subgroups, and we received expert input suggesting that there is not convincing evidence of a differential effect across subgroups. As such, we have chosen to present combined data for the PD-1 immunotherapies across these histologic subtypes.

Patients with higher levels of PD-L1 expression had better responses to all three agents, but different cutpoints and assays were used for each agent in the randomized trials (see appendix Table C5). As such, it is difficult to be certain whether the effect of PD-L1 expression is the same for the three therapies. The issue of whether histology is an effect modifier is raised by the lack of association between PD-L1 levels and response to nivolumab in patients with squamous histology in CheckMate 017. We do not have data on whether PD-L1 levels predict response to treatment in patients with squamous histology treated in POPLAR and KEYNOTE-010, however given the lack of an overall subgroup effect discussed above, we present combined data for both histologies when looking at responses at different levels of PD-L1 expression.

Although the scope of this assessment was for PD-1 immunotherapy in patients without a driver mutation, the data available to us do not provide that exact subgroup and so patients with anaplastic lymphoma kinase (ALK) and KRAS mutations are not separately broken out. We were primarily concerned about assessing therapy in patients who are EGFR-, and two of the four trials provide these subgroup data, and a third trial was in patients with squamous cell carcinoma who have a low rate of EGFR mutations.⁷ In the trial of atezolizumab, approximately 11% of patients were EGFR+ but results were not stratified. We did not detect important heterogeneity when we included this trial in the meta-analysis of EGFR- patients. However, given our estimation, as discussed below, that PD-1 immunotherapy may have no benefit in patients with EGFR+ tumors, the estimates of benefit for atezolizumab may be slightly diluted by including a percentage of patients we believe may not be receiving benefit from treatment.

PD-1 Immunotherapies for First-Line Treatment of Patients without a Driver Mutation

We currently have no direct evidence from randomized trials comparing PD-1 immunotherapies with a platinum doublet for first line treatment of advanced NSCLC. In the randomized trials, objective response rates (ORRs) were 9 to 15% with docetaxel monotherapy in patients who had received prior chemotherapy,^{16-18,70} but response rates with platinum-based doublets as first-line treatment are 24 to 30%.⁶⁷ Given this, the comparator response rate that would be expected with

standard-of-care chemotherapy should be much higher with a first-line platinum doublet than the response rate seen with second-line docetaxel. However, although we can expect that first-line PD-1 immunotherapy will be no worse than when it is used second-line, it is possible that the response rate is substantially better in this setting.

We have chosen to look at this question because patients are already being treated with first-line PD-1 immunotherapy in the absence of published evidence from randomized trials. If similar response rates to what was seen with second-line therapy are achieved with first-line PD-1 immunotherapy, it could delay or even obviate the need for chemotherapy for a proportion of patients.

Two trial results of note have been announced, but details are only currently available via press release. The Keynote 024 trial reportedly showed benefits in both OS and PFS with first-line pembrolizumab.⁷⁵ The CheckMate 026 trial reportedly showed no benefit in its primary outcome of PFS with first-line nivolumab.⁷⁶ Both of these trials were performed in patients with tumors that express PD-L1, but the cutpoints for the analyses were different and used different assays, which might explain the conflicting results. We expect these results to become available in the near future, and they will provide direct evidence on this issue.

<u>Summary</u>

There are substantial uncertainties about the efficacy of first-line PD-1 immunotherapies in NSCLC. While it seems possible that a proportion of patients would achieve durable responses that would delay the need for treatment with chemotherapy, the evidence base is insufficient ("I").

PD-1 Immunotherapies for Second- or Third-Line Treatment of EGFR+ NSCLC

We currently have no direct evidence comparing PD-1 immunotherapies with a platinum doublet as subsequent-line treatment (after TKIs) of EGFR+ advanced NSCLC. We are looking at this issue, however, as we were informed by clinical experts that some clinicians are using PD-1 immunotherapy in preference to chemotherapy in this setting.

Few EGFR+ patients were in the trials discussed above comparing PD-1 immunotherapies with docetaxel, however two such trials did report on this subgroup. As discussed above, our metaanalysis (Appendix E) suggests that the effects on OS of PD-1 immunotherapy compared with docetaxel was different in EGFR- and EGFR+ patients. This analysis suggests that there is little if any benefit with PD-1 immunotherapy compared with docetaxel in EGFR+ patients treated after progression on TKI therapy and prior to treatment with a platinum doublet. As such, there are reasons to be concerned that PD-1 immunotherapy could be inferior to a platinum doublet (which is more efficacious than docetaxel monotherapy, as discussed above).

<u>Summary</u>

Although the evidence base is insufficient ("I"), indirect evidence raises concerns that in patients with EGFR+ advanced NSCLC who have progressed after TKI therapy, treatment with PD-1 immunotherapy may be inferior to a platinum-based chemotherapy doublet. Until direct evidence is available, we feel that PD-1 immunotherapy should be avoided in this setting.

5. Other Benefits or Disadvantages

Our reviews seek to provide information on other benefits or disadvantages offered by the intervention to the individual patient, caregivers, the delivery system, other patients, or the public that would not have been considered as part of the evidence on comparative clinical effectiveness. Examples include but are not limited to:

- 1. Methods of administration that improve or diminish patient acceptability and adherence
- 2. A public health benefit, e.g., reducing new infections
- 3. Treatment outcomes that reduce disparities across various patient groups
- 4. More rapid return to work or other positive effects on productivity (if not considered a benefit as part of comparative clinical effectiveness)
- New mechanisms of action for treatments of clinical conditions for which the response to currently available treatments varies significantly among patients for unknown reasons (substantial heterogeneity of treatment effect)

As discussed above, because of the distribution of smoking behavior within the United States, lung cancer has become more common among patients in lower socioeconomic groups. Thus treatments for the disease will disproportionately affect these groups, and any economic burdens/financial toxicities of treatments will be felt more greatly. Additionally, we were told that lung cancer is a stigmatized disease, and so patients with advanced NSCLC suffer from this additional burden as well.

Although patient groups discussed with us the importance of outcomes of treatment such as reductions in distress and anxiety, these were not well captured in the clinical trials. The trials did show reductions in symptom burden with newer therapies, and it is possible that these would predict reductions in these other outcomes.

Patients with NSCLC have high rates of comorbid vascular and pulmonary disease and many such patients may not have qualified for the clinical trials that assessed these newer therapies. TKI therapy is well tolerated and can be administered even to those with a poor performance status. The safety and efficacy of PD-1 immunotherapy in such patients is less clear.

6. Comparative Value

6.1 Overview

We conducted analyses of the outcomes, costs, and cost-effectiveness of treatment for advanced NSCLC for two distinct populations:

- 1. First-line treatment with TKIs versus chemotherapy doublet (cisplatin+pemetrexed, CIS-PEM) for EGFR+ patients (population 1); and
- Second-line treatment with PD-1 immunotherapy versus docetaxel among patients without the EGFR mutation who have progressed on a first-line chemotherapy doublet (population 3).

As noted in the evidence review, there was no published or otherwise publicly-available direct comparative evidence for second- or third-line treatment with PD-1 immunotherapy in EGFR+ patients who have progressed on TKI therapy (population 2), or first-line treatment with PD-1 immunotherapy in patients without a driver mutation (population 4). Therefore, these populations were not explicitly modeled.

Analyses were carried out using a simulation model based on partition survival curves. Drug cost estimates were based on average wholesale acquisition costs and estimates of adverse events and other clinical parameters from relevant clinical trial data.

We also used outputs from this model to inform a population-based analysis of the one- and fiveyear potential budget impact of different treatment regimens. As described further in Section 6.3, we conducted analyses only for PD-1 immunotherapies, given the established and long-term presence of TKIs for first-line treatment of EGFR+ NSCLC. Potential budget impact was assessed using assumed levels of uptake over these timeframes and included assessment of drug costs as well as potential cost savings from treatment. We attempt to estimate whether the potential budget impact for any new drug at list price would surpass a threshold related to growth targets for net health care cost growth at the national level.

6.2 Incremental Costs per Outcome Achieved

Cost-Effectiveness Model: Methods

Study Aims

The primary aim of this analysis was to estimate the cost-effectiveness of various treatments for two populations. First, we compared first-line treatments for treatment-naïve patients with EGFR+

NSCLC. Second, we compared second-line treatments for NSCLC patients without the EGFR mutation who have progressed after first-line chemotherapy. The specific comparisons are given below:

- 1. First-line treatment strategies for EGFR+ NSCLC
 - CIS-PEM platinum-based chemotherapy doublet (baseline comparator)
 - TKI: Afatinib (Gilotrif[®], Boehringer Ingelheim, AFAT)
 - TKI: Erlotinib (Tarceva[®], Genentech, ERLO)
 - TKI: Gefitinib (Iressa[®], AstraZeneca, GEFI)
- 2. Second-line treatment strategies for EGFR- NSCLC
 - Docetaxel (baseline comparator, DOCX)
 - PD-1 immunotherapy: Atezolizumab (Tecentriq[®], Genentech, ATEZ)
 - PD-1 immunotherapy: Nivolumab (Opdivo[®], Bristol-Myers Squibb, NIVO)
 - PD-1 immunotherapy: Pembrolizumab (Keytruda[®], Merck, PEMB)

Key Assumptions

We made a number of key assumptions to inform our model, as described below:

Table 14. Key assumptions

Assumption	Rationale
8.9-month increase in TKI median OS vs. CIS-PEM	Comparing patients who only receive TKIs with patients who only receive chemotherapy during and after the intervention period indicates OS is approximately 9 months longer in patients who received only a TKI than in patients who received only chemotherapy
Proportional hazards assumption holds throughout	Proportional hazards modeling used in each TKI clinical trial
for TKIs	serving as input to network meta-analysis
Time-dependent HRs for immunotherapies	In recognition of PD-1 immunotherapies' violation of the proportional hazard assumption; survival tends to stabilize over time
No vial sharing occurred	Vial sharing illegal for Medicare beneficiaries receiving drugs on outpatient basis
Mean weight: 74.13 kg	Based on KEYNOTE-001 2L NSCLC cohort
Mean height: 1.78 m	To derive average patient BSA of 1.92 m2 (Burmaster 1998) ⁷⁷

OS=overall survival; CIS-PEM=cisplatin+pemetrexed; TKIs=tyrosine kinase inhibitors; HRs=hazard ratios; NSCLC=non-small cell lung cancer

Model Structure

The model framework is depicted in Figure 9. Outcomes were modeled using a partition survival approach and three health states: progression-free (PF), progression (PRO), and death (see Figure 10). Advantages of partition survival models are that they are less data intensive than other more complex modeling approaches, and that they can use commonly available data reported in clinical trial publications. For each treatment regimen, a hypothetical patient population will spend time in the PF health state and the PRO health state. The mean time, quality adjusted time, and costs spent in each health state are summed to provide estimates of life expectancy, quality adjusted life expectancy, and total costs. We used a cycle length of one week to reflect the dosing schedules for the included drug regimens. We utilized a health system perspective (i.e., direct medical care costs only) and a lifetime horizon, modeling patients from treatment initiation until death. We used a 3% discount rate for all future outcomes and costs. We developed the model in Microsoft Excel.

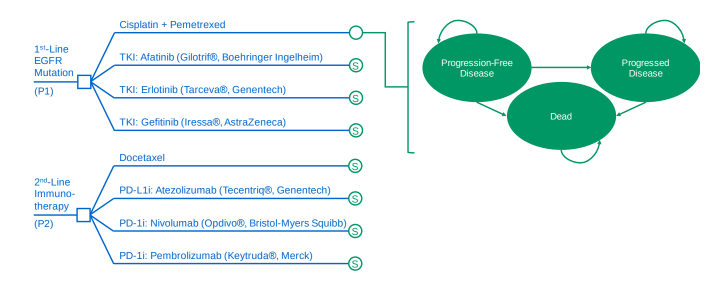
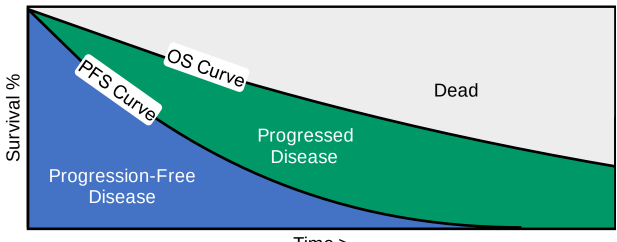


Figure 9. Model framework: Management of advanced NSCLC

Figure 10. Partition survival model approach





Treatment Strategies

The included treatment regimens were based on FDA-labeled indications for first-line EGFR+ NSCLC treatment with TKIs, and second-line NSCLC treatment with PD-1 immunotherapy among those without EGFR mutations. The primary baseline comparator in the first-line setting was a platinum-based chemotherapy doublet of cisplatin plus pemetrexed (CIS-PEM), as this as this is likely the most effective platinum doublet option for most patients with NSCLC. We recognize, however, that several recent trials have involved comparisons to other chemotherapy regimens and/or placebo. To account for the various trials and trial comparisons, a network meta-analysis was conducted (see Appendix D for further details and results).

The primary baseline comparator in the second-line setting was single-agent chemotherapy with docetaxel (DOCX). While our initial intent was also to conduct network meta-analysis to support the modeling of PD-1 immunotherapy, study populations for primary analyses were not comparable due to differences in entry criteria and/or assays used to determine PD-L1 positivity. We therefore assessed the clinical impact of each PD-1 immunotherapy agent based on treatment effects observed versus DOCX in the key clinical trial (or trials, in the case of nivolumab). The base case PD-1 immunotherapy strategies were chosen based on labeled or expected labeled indications (i.e., both PD-L1 negative and positive patients for nivolumab, PD-L1 positive only for pembrolizumab (>1%) and atezolizumab (TC 1/2/3 or IC 1/2/3).

Model Inputs: Clinical

We fit parametric survival curves to progression-free survival (PFS) Kaplan-Meier data for the universal comparators (CIS-PEM & DOCX) in both the first- and second-line settings, utilizing the

approach described by Hoyle and Henley.⁷⁸ First, we extracted data points from digitized copies of available survival curves, then used the extracted values, the number of surviving patients at each time interval, and maximum likelihood functions to estimate the underlying individual patient data. We assumed that the rate of censoring was the same between the first- and second-line settings, which allowed us to estimate the number at risk at set time points for the first- and second-line curves from the pooled number-at-risk data.

Our network meta-analysis of randomized trials showed that, for patients with EGFR+ advanced NSCLC, treatment with TKIs improves PFS compared with a platinum-based doublet, but has little effect on OS. Assessing the benefit of TKIs on OS is difficult, as there were high crossover rates in the randomized trials.^{35,37,45} As noted in Section 4, comparisons of patients who only receive TKIs with patients who only receive chemotherapy indicates OS is approximately 9 months longer in patients who received only a TKI than in those with only chemotherapy. We therefore assumed a base case in which treatment with any TKI at any point during therapy for EGFR+ advanced NSCLC improves median OS by approximately 9 months (modeled range: 6 - 13 months). A universal hazard ratio (HR 0.48; range 0.38-0.58) was derived to approximate median OS benefit, and was applied to the baseline CIS-PEM curve. For transparency, the TKI OS hazard ratios versus cisplatin-based doublets estimated by the network meta-analysis are shown in Appendix D. All base case model results reflect the use of this assumed OS benefit parameter.

Because survival data for the PD-1s are subject to a long tail and appear to violate the proportional hazards assumption, our original intent was to derive time-dependent hazard ratios using digitized survival curve data and an assumed survival distribution. However, lack of available data, particularly in the distribution's tail and for certain subgroups, prevented development of estimates with suitable precision. We therefore approximated a "flattening" of both PFS and OS for each PD-1 by assuming the full study-reported HRs for the first 10 months, followed by reductions in these HRs by 25%, 50%, and 75% for months 11-20, 21-30, and 31 and beyond. Ten-month increments were chosen because of the approximate timing of inflections in survival from the trial reports.

Model Inputs: Adverse Events

The model included Grade 3/4 adverse events that occurred in at least 5% of patients for *any* of the treatment comparators (Table 15), and were derived from key clinical trials and/or each drug's prescribing information. Because some of the adverse event rates for some regimens were reported only if the event occurred in >10% of patients, we also conducted a scenario analysis with this cutoff.

Grade 3/4 Adverse Events	CIS- PEM ³⁵	DOCX ⁷⁹	AFAT⁸ 0	ERLO ⁴ 5	GEFI ⁸¹	ATEZ ²	NIVO ⁸²	PEMB 83
Anemia	6.3%	9.0%	*	1.0%	*	*	*	5.0%
Diarrhea	0.0%	3.0%	15.0%	5.0%	3.0%	1.0%	*	0.0%
Dyspnea	*	*	*	*	*	7.0%	*	2.0%
Fatigue	12.6%	*	*	6.0%	*	*	*	7.0%
Hyponatremia	*	*	*	*	*	*	5.0%	9.0%
Infection	*	10.0%	*	*	*	*	*	*
Leukopenia	8.1%	49.0%	*	*	*	*	*	*
Nausea	3.6%	5.0%	*	*	*	1.0%	*	0.0%
Neuromotor	*	5.0%	*	*	*	*	*	*
Neutropenia	18.0%	65.0%	*	*	*	*	*	*
Paronychia/Nail disorders	0.0%	1.0%	11.0%	*	0.1%	*	*	*
Pneumonitis/Pneumonia	*	*	*	1.0%	*	6.0%	0.4%	0.2%
Pulmonary/respiratory tract	*	21.0%	*	*	*	*	*	1.0%
infection								
Rash	0.0%	*	*	13.0%	*	*	0.4%	0.0%
Skin reactions	*	1.0%	16.0%	*	2.0%	*	*	*
Stomatitis	0.9%	2.0%	9.0%	*	0.3%	*	*	*

Table 15. Adverse events per regimen

*=Not reported; CIS-PEM=cisplatin+pemetrexed; DOCX=docetaxel; AFAT=afatinib; ERLO=erlotinib; GEFI=gefitinib; ATEZ=atezolizumab; NIVO=nivolumab; PEMB=pembrolizumab

Model Inputs: Drug Utilization and Costs

The estimation of drug utilization (Appendix F) was derived from several factors, including the dosing schedule, where the dose may be fixed by weight or by body surface area (BSA; see Key Assumptions). If a regimen is based on treat-to-progression, the treatment utilization and cost were applied to all patients who remain in the PF health state over time. If a finite number of cycles is used (as with CIS-PEM), patients may remain in the PF state without active treatment. No vial sharing was assumed to occur. Drug unit costs were applied to the utilization estimates to calculate total estimated drug treatment costs.

We used the wholesale acquisition cost (WAC) for each drug and noted each available formulation (Table 16). Based on the regimen-specific dosage specified above, the model utilized the lowest-cost combination of tablets and/or vials for each regimen. For ATEZ and PEMB, we applied a one-time cost of \$274 for PD-L1 level testing in the base case analysis.

Table 16. Drug unit costs

Drug Cost Parameters	Default	< Ra	nge >	Distribution	Reference
Cisplatin per mg	\$0.36	\$0.29	\$0.43	Normal	Redbook
Cisplatin administration	\$91.72	\$73.38	\$110.06	Normal	CPT 96417 & 96415
Cisplatin dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Pemetrexed 500 mg vial	\$3,162.00	\$2,529.60	\$3,794.40	Normal	Redbook
Pemetrexed 100 mg vial	\$632.40	\$505.92	\$758.88	Normal	Redbook
Pemetrexed administration	\$136.15	\$108.92	\$163.38	Normal	CPT 96413
Pemetrexed dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Docetaxel per mg	\$9.55	\$7.64	\$11.46	Normal	Redbook
Docetaxel administration	\$136.15	\$108.92	\$163.38	Normal	CPT 96413
Docetaxel dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Afatanib 40 mg tablet	\$233.05	\$186.44	\$279.66	Normal	Redbook
Afatanib dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Erlotinib 150 mg tablet	\$241.52	\$193.22	\$289.83	Normal	Redbook
Erlotinib dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Gefitinib 250 mg tablet	\$241.20	\$192.96	\$289.44	Normal	Redbook
Gefitinib dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Atezolizumab 1200 mg vial	\$8,620.00	\$6,896.00	\$10,344.00	Normal	Redbook
Atezolizumab administration	\$136.15	\$108.92	\$163.38	Normal	CPT 96413
Atezolizumab dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Nivolumab 100 mg vial	\$2,470.48	\$1,976.38	\$2,964.58	Normal	Redbook
Nivolumab 40 mg vial	\$988.19	\$790.55	\$1,185.83	Normal	Redbook
Nivolumab administration	\$136.15	\$108.92	\$163.38	Normal	CPT 96413
Nivolumab dose intensity	100.0%	80.0%	100.0%	Beta	Assumption
Pembrolizumab 100 mg vial	\$4,380.74	\$3,504.59	\$5,256.89	Normal	Redbook
Pembrolizumab administration	\$136.15	\$108.92	\$163.38	Normal	CPT 96413
Pembrolizumab dose intensity	100.0%	80.0%	100.0%	Beta	Assumption

Costs per adverse event were based on data from the Centers for Medicare and Medicaid Services (CMS) list of Medicare Severity-Diagnosis Related Groups (MS-DRGs) for the fiscal year 2016,⁸⁴ and are shown in Appendix F.

To estimate costs in the progression health state, we assumed the following subsequent therapies for the modeled populations: (1) first-line TKI-treated patients received a chemotherapy doublet of CIS-PEM; (2) first-line CIS-PEM patients received DOCX; (3) second-line PD-1 immunotherapy patients received DOCX; and (4) second-line DOCX patients received gemcitabine monotherapy. The cost of each subsequent regimen was multiplied by the proportion of patients entering the progressed disease health state during each weekly model cycle. Subsequent regimen costs were derived by calculating the average weekly cost of survival-linked regimens for DOCX and gemcitabine (GEM)⁸⁵ over 3 months for third-line and CIS-PEM over 12 months for second-line

treatment (Table 17). DOCX and CIS-PEM survival for calculating post-progression weekly cost was assumed to be equivalent to the baseline curves in the two models.

Original	Subsequent	Cost/
Treatment	Treatment	Week
CIS-PEM	DOCX	\$441
DOCX	GEM	\$82
AFAT	CIS-PEM	\$605
ERLO	CIS-PEM	\$605
GEFI	CIS-PEM	\$605
ATEZ	DOCX	\$441
NIVO	DOCX	\$441
PEMB	DOCX	\$441

Table 17. Post-progression costs

CIS-PEM=cisplatin+pemetrexed; DOCX=docetaxel;

AFAT=afatinib; ERLO=erlotinib; GEFI=gefitinib;

ATEZ=atezolizumab; NIVO=nivolumab;

PEMB=pembrolizumab; GEM=gemcitabine

Model Inputs: Health State Utilities

Health state utilities were derived from publicly available literature and applied to the disease states of progression-free and progressed disease (Appendix F). We assumed that health state utility values did not vary across the treatments evaluated in the model. For the progression-free health state, different utilities were applied depending on whether the patient was receiving first-or second-line treatment, to represent decreased quality of life due to progression following first-line treatment. We applied a regimen-weighted disutility for experiencing any Grade 3/4 adverse event; the total percentage of patients who experienced any Grade 3/4 adverse events for each regimen was multiplied by the adverse event disutility and then subtracted from the first month of PFS for each regimen. We assumed that the total time with a Grade 3/4 adverse event for patients experiencing any Grade 3/4 adverse event was one month.

Model Outcomes

The model estimated the amount of time, on average, patients spend progression-free and in progression. Unadjusted and utility-adjusted time spent in each health state was summed to provide estimates of life expectancy and quality-adjusted life expectancy.

Model outcomes of interest for each intervention included:

• Quality adjusted life expectancy (discounted)

- Life expectancy (discounted)
- Mean time in the progression-free and post-progression health states (discounted)
- Pre-progression, post-progression, and total costs (discounted)
- Incremental cost-effectiveness ratios (ICER) for each intervention versus the standard comparator (CIS-PEM or DOCX), in pairwise comparisons

Sensitivity Analyses

The model programming allows for flexible and comprehensive sensitivity analyses. One-way sensitivity analyses used 95% confidence intervals from clinical evidence as ranges, where available. When 95% confidence intervals were not available, uncertainty ranges were based on plausible values from the published literature. We also conducted a probabilistic sensitivity analysis (PSA) by jointly varying all model parameters over 4,000 simulations, then calculating 95% credible range estimates for each model outcome.

Finally, we ran seven scenario analyses: (1) omitting our assumption of an 8.9-month overall survival benefit and utilizing NMA OS hazard ratios from TKI/CIS-PEM crossover populations; (2 & 3) using ATEZ PFS and OS hazards ratios derived from POPLAR intention to treat (ITT) and high PD-L1 expression on TC3 or IC3 populations; (4 & 5) using NIVO PFS and OS hazard ratios derived from CheckMate 017/057 populations with PD-L1 levels of >1% and >10%; (6) using PEMB PFS and OS hazard ratios derived from KEYNOTE-010 populations with PD-L1 levels of >50%; and (7) in recognition of different thresholds for adverse event reporting among study drugs, we explored a scenario in which we only included adverse events with at least one drug reporting >10% of patients experiencing the event instead of the base case threshold of >5%.

Cost-Effectiveness Model: Results

Base Case Results

The results of the pairwise comparisons are provided in Table 18a and Table 18b for the first-line setting with TKIs, and Table 19a and Table 19b for the second-line setting with PD-1 immunotherapies. These tables report detailed results for each regimen in each line as well as the incremental results versus their respective baseline comparators. Note that survival results in the table will not match those seen in clinical trials because of our anchoring of hazard ratios to the baseline survival curves for CIS-PEM and DOCX (rather than use of observed survival curves in each trial).

All results presented below are based on deterministic analyses (i.e., based on point estimates for all model parameters), due to wide confidence intervals for some of the model estimates. Results from companion probabilistic analyses (which assume a distribution for each parameter) can be found in Appendix F

Table 18a. Results by regimen for first-line EGFR+ patients

	CIS-PEM	AFAT	ERLO	GEFI
Total Costs	\$112,361	\$194,046	\$192,519	\$178,175
Drug Costs	\$32,042	\$91,560	\$91,463	\$76,609
PFS Supp. Care Costs	\$10,217	\$20,757	\$19,988	\$16,696
Administration Costs	\$1,145	\$0	\$0	\$0
Progression Costs	\$15,763	\$31,063	\$32,290	\$37,573
Death Costs	\$48,192	\$46,923	\$46,923	\$46,923
Adverse Event Costs	\$5,002	\$3,744	\$1,855	\$375
Total QALYs	0.88	1.50	1.50	1.48
PFS QALYs	0.42	0.84	0.81	0.68
Progression QALYs	0.46	0.66	0.68	0.80
Total Life Years (OS)	1.22	2.06	2.06	2.06
PFS LYs	0.54	1.08	1.04	0.87
Progression LYs	0.68	0.98	1.02	1.19
Median PFS (months)	5.1	10.6	10.2	8.5
Median OS (months)	12.5	21.4	21.4	21.4

CIS-PEM=cisplatin+pemetrexed; DOCX=docetaxel; AFAT=afatinib; ERLO=erlotinib; GEFI=gefitinib; PFS=progression-free survival; OS=overall survival; QALY=quality-adjusted life year

Table 18b. Incremental results for first-line EGFR+ patients

Incremental Results				
	CIS-PEM	AFAT	ERLO	GEFI
ICER		\$131,051	\$129,497	\$109,666
Incremental Costs		\$81,685	\$80,158	\$65,814
Drug Costs		\$59,519	\$59,421	\$44,567
PFS Supp. Care Costs		\$10,540	\$9,771	\$6,479
Administration Costs		-\$1,145	-\$1,145	-\$1,145
Progression Costs		\$15,299	\$16,527	\$21,810
Death Costs		-\$1,269	-\$1,269	-\$1,269
Adverse Event Costs		-\$1,259	-\$3,147	-\$4,628
Incremental QALYs		0.62	0.62	0.60
PFS QALYs		0.42	0.39	0.26
Progression QALYs		0.20	0.23	0.34
Incremental Life Years (OS)		0.84	0.84	0.84
PFS LYs		0.54	0.50	0.33
Progression LYs		0.30	0.34	0.50

CIS-PEM=cisplatin+pemetrexed; DOCX=docetaxel; AFAT=afatinib; ERLO=erlotinib; GEFI=gefitinib; ICER=incremental cost effectiveness ratio; PFS=progression-free survival; OS=overall survival; QALY=quality-adjusted life year

Use of each of the first-line TKI regimens (including the assumption of an 8.9-month gain in median survival for each regimen) resulted in a 0.84 life-year gain in survival relative to CIS-PEM. On a quality-adjusted basis, QALYs gained versus CIS-PEM were also very similar, ranging from 0.60 for GEFI to 0.62 for AFAT and ERLO. Incremental costs versus CIS-PEM were lower for GEFI (~\$66,000) than for the other TKIs, as a function of a shorter duration of time spent in the progression-free state (and a consequently shorter duration of treatment). Cost-effectiveness estimates were similar across the TKIs, ranging from approximately \$110,000 - \$130,000 per QALY gained.

Table 19a. Results by regimen for second-line PD-1 immunotherapy patients

	DOCX	ATEZ: TC1/2/3 or IC1/2/3	NIVO: All Comers	PEMB: PD-L1 >1%
Total Costs	\$87,831	\$170,455	\$165,802	\$154,051
Drug Costs	\$9,563	\$84,641	\$83,929	\$72,760
PFS Supp. Care Costs	\$7,224	\$10,423	\$10,204	\$8,704
Administration Costs	\$947	\$1,333	\$1,927	\$1,126
Progression Costs	\$2,134	\$25,696	\$21,571	\$21,764
Death Costs	\$48,693	\$47,493	\$47,804	\$47,912
Adverse Event Costs	\$19,270	\$870	\$367	\$1,785
Total QALYs	0.48	0.88	0.79	0.74
PFS QALYs	0.24	0.36	0.35	0.30
Progression QALYs	0.24	0.52	0.44	0.44
Total Life Years (OS)	0.88	1.66	1.47	1.40
PFS LYs	0.38	0.54	0.53	0.46
Progression LYs	0.50	1.12	0.94	0.95
Median PFS (months)	3.0	3.7	4.2	3.5
Median OS (months)	8.8	14.1	12.5	11.8

ATEZ=atezolizumab; TC1/2/3 or IC1/2/3=PD-L1 expression ≥1%; NIVO=nivolumab; PEMB=pembrolizumab; PFS=progression-free survival; OS=overall survival; QALY=quality-adjusted life year;

Table 19b. Incremental results for second-line PD-1 immunotherapy patients

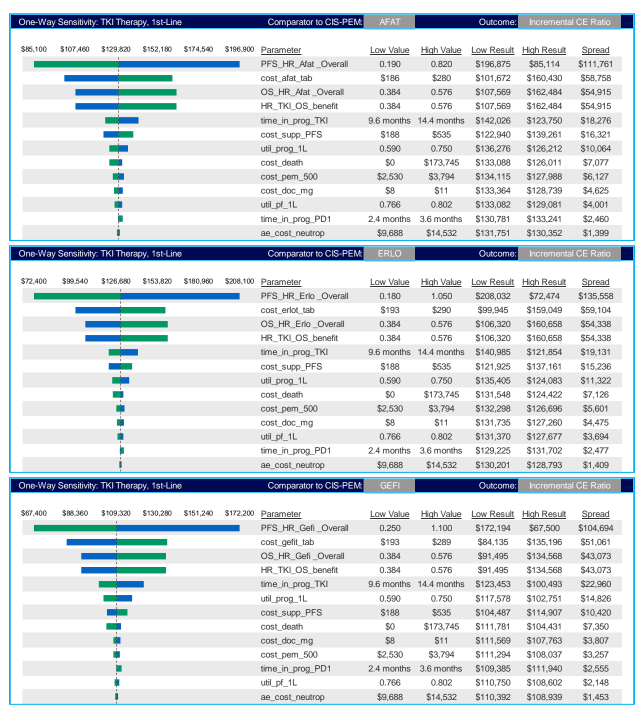
	DOCX	ATEZ: TC1/2/3 or IC1/2/3	NIVO: All Comers	PEMB: PD-L1 >1%
ICER		\$208,144	\$254,624	\$254,776
Incremental Costs		\$82,624	\$77,971	\$66,220
Drug Costs		\$75,078	\$74,365	\$63,197
PFS Supp. Care Costs		\$3,199	\$2,980	\$1,480
Administration Costs		\$386	\$980	\$179
Progression Costs		\$23,562	\$19,437	\$19,630
Death Costs		-\$1,201	-\$889	-\$781
Adverse Event Costs		-\$18,400	-\$18,902	-\$17,485
Incremental QALYs		0.40	0.31	0.26
PFS QALYs		0.12	0.10	0.05
Progression QALYs		0.29	0.20	0.21
Incremental Life Years (OS)		0.78	0.59	0.52
PFS LYs		0.16	0.15	0.08
Progression LYs		0.61	0.44	0.44

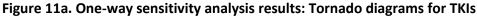
ATEZ=atezolizumab; TC1/2/3 or IC1/2/3=PD-L1 expression ≥1%; NIVO=nivolumab; PEMB=pembrolizumab; ICER=incremental cost effectiveness ratio; PFS=progression-free survival; OS=overall survival; QALY=quality-adjusted life year

Use of each of the second-line immunotherapy regimens resulted in a gain in survival (range: 0.52 for PEMB to 0.78 for ATEZ) relative to DOCX. On a quality-adjusted basis, QALYs gained versus CIS-PEM ranged from 0.26 for PEMB to 0.40 for ATEZ. Incremental costs versus DOCX ranged from a low of approximately \$66,000 for PEMB to approximately \$82,000 for ATEZ. Cost-effectiveness estimates ranged from approximately \$208,000 per QALY gained relative to DOCX for ATEZ to ~\$250,000 per QALY for the other PD-1s. Again, it is important to stress that this analysis was based on the within-trial experience for each agent. The contrast of primary interest is on the incremental outcomes, costs, and cost-effectiveness of each PD-1 in relation to DOCX, not on comparisons between the PD-1s themselves.

Sensitivity Analyses

Detailed findings from the one-way sensitivity analyses can be found below. In each one-way analysis, results were most sensitive to PFS and OS hazard ratios, drug costs, and (for TKIs) the assumption of an 8.9-year OS benefit.





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One-Way	Sensitivity	: Immunoth	nerapy, 2nd	-Line		Comparator to DOCX:	ATEZ: TC1/2	2/3 or IC1/2/3	Outcome:	Incrementa	I CE Ratio
\$135,500	\$191,240	\$246,980	\$302,720	\$358,460	\$414,200	Parameter	Low Value	High Value	Low Result	High Result	Spread
¢ 100,000	¢101,210	\$210,000	\$002,i 20	4000,100	¢111,200	OS_HR_Atez	0.400	0.850	\$135,571	\$414,122	\$278,551
						PFS_HR_Atez	0.630	1.160	\$267,319	\$156,045	\$111,273
						cost_atezo_1200	\$6,896	\$10,344	\$165,637	\$250,651	\$85,015
						util_prog_2L	0.430	0.520	\$221,877	\$193,197	\$28,680
						cost_doc_mg	\$8	\$11	\$201,182	\$215,106	\$13,924
	-					cost_death	\$0 \$0	\$173,745	\$211,169	\$200,659	\$10,509
	- i					ae_cost_neutrop	\$9,688	\$14,532	\$212,110	\$204,178	\$7,932
						cost_supp_PFS	\$188	\$535	\$204,279	\$212,056	\$7,778
						util_pf_2L	0.610	0.700	\$211,618	\$203,959	\$7,658
						ae_cost_leukop	\$9,688	\$14,532	\$211,134	\$205,155	\$5,979
						time_in_prog_PD1	2.4 months	3.6 months	\$209,323	\$203,547	\$5,776
	- 1					ae_cost_respinfx	\$8,251	\$12,377	\$209,236	\$207,053	\$2,183
						cost_doc_admin	\$109	\$163	\$207,455	\$208,834	\$1,379
	0			1.5							
ne-way	Sensitivity	: immunotr	ierapy, zno	-Line		Comparator to DOCX:	NIVO: AI	l Comers	Outcome:	Incrementa	I CE Ratio
\$181,700	\$222,240	\$262,780	\$303,320	\$343,860	\$384,400	Parameter	Low Value	High Value	Low Result	High Result	Spread
						OS_HR_Nivo	0.551	0.825	\$189,467	\$384,342	\$194,875
						PFS_HR_Nivo	0.520	1.126	\$345,095	\$181,728	\$163,368
						cost_nivol_100	\$1,976	\$2,965	\$208,944	\$300,303	\$91,359
						util_prog_2L	0.430	0.520	\$269,967	\$237,734	\$32,233
						vial_sh_nivol	Yes	No	\$234,513	\$254,624	\$20,110
						cost_nivol_40	\$791	\$1,186	\$245,488	\$263,760	\$18,272
		-				cost_doc_mg	\$8	\$11	\$248,051	\$261,197	\$13,146
						util_pf_2L	0.610	0.700	\$259,806	\$248,429	\$11,377
						ae_cost_neutrop	\$9,688	\$14,532	\$259,765	\$249,483	\$10,282
		-				cost_death	\$0	\$173,745	\$257,526	\$247,440	\$10,087
		-				cost_supp_PFS	\$188	\$535	\$249,956	\$259,348	\$9,393
						ae_cost_leukop	\$9,688	\$14,532	\$258,499	\$250,748	\$7,751
						time_in_prog_PD1	2.4 months	3.6 months	\$256,008	\$249,831	\$6,178
ne-Way	Sensitivity	: Immunoth	nerapy, 2nd	-Line		Comparator to DOCX:	PEMB:P	D-L1 >1%	Outcome:	Incrementa	I CE Ratio
\$183,800	\$232,940	\$282,080	\$331,220	\$380,360	\$429,500	Parameter	Low Value	High Value	Low Result	High Result	Spread
						OS HR Pemb	0.580	0.880	\$183,836	\$429,459	\$245,623
						cost pembro 100	\$3,505	\$5,257	\$198,999	\$310,553	\$111,554
			-			PFS_HR_Pemb	0.740	1.050	\$291,666	\$220,819	\$70,847
						util_prog_2L	0.430	0.520	\$273,439	\$234,749	\$38,691
						cost doc mg	\$8	\$11	\$246,897	\$262,656	\$15,759
						ae_cost_neutrop	\$9,688	\$14,532	\$260,833	\$248,719	\$12,114
						cost_death	\$0	\$173,745	\$257,781	\$247,340	\$10,440
						ae_cost_leukop	\$9,688	\$14,532	\$259,342	\$250,210	\$9,132
						time_in_prog_PD1	2.4 months	3.6 months	\$256,416	\$249,065	\$7,351
		1				util_pf_2L	0.610	0.700	\$257,785	\$251,113	\$6,671
						cost_supp_PFS	\$188	\$535	\$252,046	\$257,540	\$5,494
	1										
						vial sh pembro	Yes	No	\$249,924	\$254,776	\$4,853

Figure 11b. One-way sensitivity analysis results: Tornado diagrams for immunotherapies

Results of our probabilistic sensitivity analysis (PSA) can be found in Appendix F. Our findings show substantial variability in model outcomes, particularly in second-line immunotherapies. However, the range of possible incremental cost-effectiveness ratios approached commonly-cited thresholds

(i.e., \$50,000 - \$150,000 per QALY gained) for first-line TKI regimens. In contrast, the possible range of results from PSAs on second-line immunotherapies are substantially wide due to parameter uncertainty, particularly in PFS and OS hazard ratios; thus, mean ICERs should be interpreted with great caution.

Scenario analysis results are presented in Appendix F. In general, the scenario results that involve adjustments to PFS and OS based on specific populations should be interpreted with great caution, as the hazard ratios used to derive survival were highly uncertain in most cases. A full accounting of parameter uncertainty in PSA would lead to notably different results in many cases. With this caveat, we present the deterministic estimates to explore the hypothetical impacts of targeted NSCLC therapy.

In scenario 1, we omitted our assumption of an 8.9-month overall survival benefit, and utilized network meta-analysis OS hazard ratios from TKI/CIS-PEM crossover populations. As expected, this resulted in a notable decrease in overall survival for TKIs, which lowered both drug costs and progression costs because fewer patients were alive to accrue these costs. However, clinical benefits were also severely diminished, leading to inferior survival and QALY estimates relative to CIS-PEM in some circumstances, and high cost-effectiveness ratios in others.

Scenarios 2 & 3 utilized ATEZ PFS and OS hazard ratios derived from POPLAR intention to treat (ITT) and high PD-L1 expression TC3 or IC3 populations, respectively. Survival in the ITT population was comparatively worse than the base case population, resulting in decreased survival and decreased cost but a somewhat higher cost-effectiveness ratio. In contrast, the high PD-L1 expression TC3 or IC3 population exhibited greater survival, particularly after the first 6 months of therapy, resulting in a notable increase in both overall and quality-adjusted survival. However, this also substantially increased ATEZ drug cost due to a more pronounced progression-free survival benefit than that seen in the base case.

We used NIVO PFS and OS hazard ratios derived from the CheckMate 017/057 populations with PD-L1 levels of >1% and >10% in scenarios 4 & 5. When limiting our analysis to patients with observed increased PD-L1 level versus the NIVO base case including all comers, overall and quality-adjusted survival as well as cost increased, but decreased the NIVO cost-effectiveness ratio relative to the base case in both situations.

In Scenario 6 using PEMB PFS and OS hazard ratios derived from the KEYNOTE-010 population with PD-L1 levels of >50%, we observed a near doubling of overall and quality-adjusted survival. While time in the progression-free state and associated drug costs also increased, the cost-effectiveness ratio in this scenario was reduced relative to the base case.

Finally, the scenario in which we included AEs with at least one drug reporting >10% of patients experiencing the event instead of the base case threshold of >5% showed only a modest decrease in cost for all drug regimens, and did not have a notable impact on overall results.

6.3 Potential Budget Impact

We also used the cost-effectiveness model to estimate the potential total budgetary impact of PD-1 immunotherapy treatments for NSCLC patients, based on assumed patterns of product uptake.

Potential Budget Impact Model: Methods

We used results from the same model employed for the cost-effectiveness analyses to estimate total potential budget impact. Potential budget impact was defined as the total incremental cost of using PD-1 immunotherapy rather than docetaxel for the treated population, calculated as incremental health care costs (including drug costs) minus any offsets in these costs from averted health care events. We did not include the other population modeled above (adults with advanced NSCLC who have an EGFR+ tumor and have not previously been treated for advanced disease) in this budget impact analysis, as all three of the TKIs evaluated in that population are in established use based on clinical guidelines, and with many years of market experience in two of the three cases. We also do not consider the potential budgetary implications of using these drugs for other indications than NSCLC.

Note that this analysis is performed from an *ex ante* perspective; that is, it treats all of the drugs being evaluated as though they will be new to market, whether or not they have already been launched. We did allow for differential uptake of the PD-1s by product, however, based on currently-available market share data. We estimated the net costs of using each drug rather than docetaxel, assuming no current use of the drug. All costs were undiscounted and estimated over one- and five-year time horizons. The five-year timeframe was of primary interest, given the potential for cost offsets to accrue over time.

The potential budget impact analysis included the entire candidate population for treatment, which was considered to be adults with advanced NSCLC who have a tumor that has progressed after first-line treatment with a platinum-based chemotherapy doublet.

To estimate the size of the potential candidate population for treatment with atezolizumab, we first determined the estimated prevalence of NSCLC in the US. Lung cancer prevalence in 2013 was estimated to be 415,707 patients.⁸⁶ Of those, 85% are estimated to have NSCLC and 70% of those to have advanced disease.⁸⁷ This would result in an estimate of 247,346 persons with advanced NSCLC in the US. Pembrolizumab is approved for use in second-line treatment in patients whose tumors demonstrate PD-L1 expression, and we anticipate that atezolizomab will be approved for a similar

population. Taube has reported that approximately 60% of NSCLC cancer specimens demonstrated PD-L1 expression. Stinchcombe and Socinski state that: "While it is difficult to estimate the proportion of patients who receive second-line treatment, approximately 40%–50% of patients did so in recent first-line trials."⁸⁸ We therefore assumed that 40% of patients would receive second-line treatment. Applying these percentages resulted in a candidate population size of approximately 59,400 individuals in the US. Use of nivolumab is not restricted to use in patients whose tumors express PD-L1. We therefore applied the 40% assumption for second-line treatment to the 40% of patients with tumors that do not express PD-L1, resulting in an additional candidate population for nivolumab of approximately 39,600 patients (i.e., a total of 99,000 individuals).

ICER's methods for estimating potential budget impact and calculating value-based benchmark prices are described in detail <u>elsewhere</u>.⁸⁹ Briefly, our calculations assume that the utilization of new drugs occurs without any payer, provider group, or pharmacy benefit management controls in place, to provide an estimate of "unmanaged" drug uptake by five years after launch.

In general, we examine six characteristics of the drug or device and the marketplace to estimate "unmanaged" uptake. These characteristics are listed below:

- Magnitude of improvement in clinical safety and/or effectiveness
- Patient-level burden of illness
- Patient preference (ease of administration)
- Proportion of eligible patients currently being treated
- Primary care versus specialty clinician prescribing/use
- Presence or emergence of competing treatments of equal or superior effectiveness

Based on our assessment of these criteria, we assign a new drug or device to one of four categories of unmanaged drug uptake patterns: 1) very high (75% uptake by year 5); 2) high (50% uptake by year 5); 3) intermediate (25% uptake by year 5); and 4) low (10% uptake by year 5). In this analysis, we assumed a 50% uptake pattern for nivolumab, and 25% uptake each for atezolizumab and pembrolizumab in the eligible PD-L1 positive population. We assumed that uptake would be high for nivolumab because it does not require PD-L1 testing, and intermediate for atezolizumab and pembrolizumab because of the need for PD-L1 testing in second-line NSCLC patients. This is in line with recent market share information for nivolumab and pembrolizumab.^{90,91} For the eligible population of second-line NSCLC patients whose tumors do not express PD-L1, we assumed a 75% uptake pattern for nivolumab. We assumed that uptake would be very high in this population because of the lack of other effective treatment alternatives.

Using this approach to estimate potential budget impact, we then compared our estimates to a budget impact threshold that represents a potential trigger for policy mechanisms to improve affordability, such as changes to pricing, payment, or patient eligibility. As described in <u>ICER's</u>

methods presentation, this threshold is based on an underlying assumption that health care costs should not grow much faster than growth in the overall national economy. From this foundational assumption, our potential budget impact threshold is derived using an estimate of growth in US gross domestic product (GDP) +1%, the average number of new drug approvals by the FDA each year, and the contribution of spending on retail and facility-based drugs to total health care spending. Calculations are performed as shown in Table 20.

For 2015-16, therefore, the five-year annualized potential budget impact threshold that should trigger policy actions to manage affordability is calculated to total approximately \$904 million per year for new drugs.

Item	Parameter	Estimate	Source
1	Growth in US GDP, 2015-2016 (est.) +1%	3.75%	World Bank, 2015
2	Total health care spending (\$)	\$3.08 trillion	CMS NHE, 2014
3	Contribution of drug spending to total health care spending (%)	13.3%	CMS National Health Expenditures (NHE), Altarum Institute, 2014
4	Contribution of drug spending to total health care spending (\$) (Row 2 x Row 3)	\$410 billion	Calculation
5	Annual threshold for net health care cost growth for ALL new drugs (Row 1 x Row 4)	\$15.4 billion	Calculation
6	Average annual number of new molecular entity approvals, 2013-2014	34	FDA, 2014
7	Annual threshold for average cost growth per individual new molecular entity (Row 5 ÷ Row 6)	\$452 million	Calculation
8	Annual threshold for estimated potential budget impact for each individual new molecular entity (doubling of Row 7)	\$904 million	Calculation

Table 20. Calculation of potential budget impact threshold

Potential Budget Impact Model: Results

Table 21 below presents the potential budget impact of one year and five years of utilization of each drug in the candidate population, assuming the uptake patterns previously described. Results are presented for both one-year and five-year time horizons.

Results from the potential budget impact model showed that in the first year, with the uptake pattern assumptions mentioned above, atezolizumab and pembrolizumab would each be given to an estimated 2,970 individuals, and nivolumab to approximately 11,880 (5,940 each from the PD-L1 positive and negative populations). Over the entire five-year time horizon, we estimate that

"unmanaged" uptake would lead to approximately 14,850 persons receiving atezolizumab, 14,850 receiving pembrolizumab, and 59,400 receiving nivolumab for one or more years.

After one year of treatment, net annual costs ranged from \$35,300 per patient for atezolizumab to \$42,200 per patient for nivolumab. One-year budget impact is estimated to be \$104.6 million for atezolizumab, \$112.7 million for pembrolizumab, and \$500.9 million for nivolumab. Total budget impact for nivolumab is much higher due to the larger number of patients assumed to receive nivolumab.

Across the full five-year time horizon, the weighted potential budgetary impact (i.e., adjusted for differing periods of drug utilization and associated cost-offsets) ranges from approximately \$55,500 per patient receiving pembrolizumab to \$64,600 per patient receiving nivolumab. Average budget impact per year is estimated as approximately \$169.4 million for atezolizumab, \$164.6 million for pembrolizumab, and \$766.5 million for nivolumab. This annualized potential budget impact is 19% of the budget impact threshold of \$904 million for atezolizumab, 18% of the threshold for pembrolizumab, and 85% of the threshold for nivolumab.

		Analytic Horizon = 1 Year			Analytic Horizon = 5 Years			
	Eligible	Number	Annual BI per	Total BI	Number	Weighted BI	Average BI	
	Population	Treated	Patient*	(millions)	Treated	per Patient*	per year	
							(millions)	
Atezolizumab	59,400	2,970	\$35,300	\$104.6	14,850	\$57,100	\$169.4	
Pembrolizumab	59,400	2,970	\$38,000	\$112.7	14,850	\$55,500	\$164.6	
Nivolumab ⁺	99,000	11,880	\$42,200	\$500.9	59,400	\$64,600	\$766.5	

Table 21. Estimated total potential budget impact (BI) of atezolizumab

*Weighted budget impact calculated by subtracting cost offsets from drug costs for one-year horizon. For fiveyear horizon, drug costs and cost offsets apportioned assuming 20% of patients in uptake target initiate therapy each year. Those initiating in year 1 receive full drug costs and cost offsets, those initiating in year 2 receive 80% of drug costs and cost offsets, etc.

+Includes PD-L1 positive and negative patients.

6.4 Draft Value-based Benchmark Prices

Value-based price benchmarks will be provided as part of the full Evidence Report.

6.5 Summary and Comment

The primary aim of this analysis was to estimate the cost-effectiveness of treating NSCLC patients with first-line TKIs versus a chemotherapy doublet (CIS-PEM) for EGFR+ patients (population 1), and second-line treatment with PD-1 immunotherapy versus DOCX among patients who have progressed on a first-line chemotherapy doublet (population 3). For first-line treatment using TKIs targeted at an EGFR mutation, our primary analysis generated incremental cost-effectiveness ratios

ranging between approximately \$110,000 and \$130,000 per QALY gained relative to comparator treatment with CIS-PEM. These ratios are within commonly-cited thresholds for the costeffectiveness of health interventions (i.e., \$50,000-\$150,000 per QALY gained), although both deterministic and probabilistic sensitivity analyses suggest some uncertainty in these findings. There is additional uncertainty created by our need to estimate OS benefits from observational data, given the effects of cross-overs in the randomized trials. Our results for second-line PD-1 immunotherapies were more uncertain. Cost-effectiveness ratios ranged between \$208,000 and \$250,000 per QALY gained in base case analyses, and ranged similarly in scenario analyses based on levels of PD-L1 positivity due to tradeoffs between improved survival and higher drug costs as a result of longer time to progression. However, findings for all analyses varied widely in both deterministic and probabilistic sensitivity analyses as a result of wide confidence regions around key parameters such as PFS and OS hazard ratios. These results should therefore be interpreted with caution.

We note several additional limitations of our analysis. The cost-effectiveness analysis was conducted from a health system perspective, and so does not incorporate costs and effects that might be relevant from a societal perspective, such as productivity, transportation, or caregiver costs. However, the largest cost driver and a highly sensitive parameter in our model was the costs of the drugs themselves, and all patients were assumed to have a similar severity of disease. Any residual differences in transportation time or time in treatment would be unlikely to have materially affected our findings. We also assumed that there would be no vial sharing for any infused drug, in the absence of published and credible data on the frequency of this practice in NSCLC. If vial sharing does occur in actual practice for some patients, our analysis would overestimate drug costs for the affected regimens, although to a currently unknown extent.

In addition, our assumptions regarding treatment received after progression necessitated assumptions that do not reflect current clinical practice. For example, patients with NSCLC treated with chemotherapy typically receive maintenance therapy (e.g., pemetrexed), and many of those who progress on first-line TKI treatment now receive the TKI osimertinib as second-line therapy. However, most of the original trials of the TKIs were performed before these were standard options, and consideration of these regimens would require additional assumptions regarding survival after first-line progression in the absence of adequate data.

While our analysis included reported adverse events that occurred in at least 5% of patients for any regimen of interest, we did exclude adverse events that occurred in <5% of patients across all regimens, which may have ruled out certain rare but expensive events. However, given that drug costs represented the majority of total costs for any given regimen in our analysis, the effects of adding rare adverse events to our analysis would not have materially changed our findings.

Finally, the levels of regimen uptake in the marketplace by five years were based on reasoned assumptions regarding current market share and likely uptake, but actual uptake and market share may vary from these estimates. Additionally, costs for drugs already on the market (i.e., PEMB) were not considered as part of the background treatment costs; rather, the potential budget impact analysis was performed from the perspective of replacing the comparator regimen with each of the PD-1s.

In summary, targeted regimens for first- and second-line use in NSCLC appears to confer clinical benefits in terms of lengthening progression-free and overall survival as well as improved quality of life. At current wholesale acquisition costs, the estimated cost-effectiveness of each of the TKIs appears to fall within commonly-accepted thresholds. While the cost-effectiveness of PD-1 immunotherapies exceeds these thresholds, there is greater uncertainty in these findings given variability in estimates of overall and progression-free survival.

This is the first Midwest CEPAC review of treatment options for non-small cell lung cancer.

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APPENDICES

- A. Evidence Review Methods
- B. PRISMA and Evidence Review Table
- C. Additional Results from Evidence Review
- D. Network Meta-Analysis Methods and Results
- E. Subgroup Meta-Analysis Methods and Results
- F. Comparative Value Supplemental Information
- G. Previous Technology Assessments and Systematic Reviews
- H. Ongoing Studies

Appendix A. Evidence Review Methods

Table A1. PRISMA 2009 Checklist

		TITLE
Title	1	Identify the report as a systematic review, meta-analysis, or both.
		ABSTRACT
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number. INTRODUCTION
Rationale	3	Describe the rationale for the review in the context of what is already known.
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).
		METHODS
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I ²) for each meta-analysis.
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.
		RESULTS
Study	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for
selection Study	18	exclusions at each stage, ideally with a flow diagram. For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up
characteristics	10	period) and provide the citations.

		RESULTS (continued)
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).
		DISCUSSION
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.
		FUNDING
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.

Moher D, Liberati A, Tetzlaff J, Altman DG. The PRISMA Group (2009). Preferred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. *PLoS Med* 6(6): e1000097. doi:10.1371/journal.pmed1000097

1	Erlotinib.ti,ab
2	Gefitinib.ti,ab
3	Afatinib.ti,ab
4	Nivolumab.ti,ab
5	Pembrolizumab.ti,ab
6	Atezolizumab.ti,ab
7	1 or 2 or 3 or 4 or 5 or 6
8	randomized controlled trial.pt.
9	controlled clinical trial.pt.
10	randomized.ab.
11	placebo.ab.
12	drug therapy.fs.
13	randomly.ab.
14	trial.ab.
15	groups.ab.
16	observational study.pt.
17	exp case-control studies/
18	exp cohort studies/
19	exp cross-over studies/
20	exp matched-pair analysis/
21	multicenter study.pt.
22	8 or 9 or 10 or 11 or 12 or 13 or 14 or 15
23	16 or 17 or 18 or 19 or 20 or 21
24	comparative study.pt. or compare.ab,ti. or compares.ab,ti. or compared.ab,ti. or comparing.ab,ti. or
	comparison.ab,ti. or comparison.ab,ti. or comparative.ab,ti. or effective.ab,ti. or effectiveness.ab,ti. or
	versus.ab,ti. or vs.ab,ti.
25	23 and 24
26	22 or 25
27	exp carcinoma, non-small-cell lung/
28	(lung and (cancer\$ or carcin\$ or neoplasm\$ or tumour\$ or tumor\$) and ((non-small or nonsmall) and
20	cell)).ti,ab.
29	nsclc.ti,ab.
30	27 or 28 or 29
31	7 and 26 and 30
32	exp animals/
33	humans.sh.
34	32 not 33
35	31 not 34
36	limit 35 to english language
37	(case reports or comment or congresses or editorial or letter or review).pt
38	36 not 37

Table A2. Search strategy of Ovid Medline on June 8, 2016

Table A3. Search strategy of Cochrane Central Register of Controlled Trials on June 8, 2016

1	
1	Erlotinib.ti,ab
2	Gefitinib.ti,ab
3	Afatinib.ti,ab
4	Nivolumab.ti,ab
5	Pembrolizumab.ti,ab
6	Atezolizumab.ti,ab
7	1 or 2 or 3 or 4 or 5 or 6
8	exp carcinoma, non-small-cell lung/
9	(lung and (cancer\$ or carcin\$ or neoplasm\$ or tumour\$ or tumor\$) and ((non-small or nonsmall) and
	cell)).ti,ab.
10	nsclc.ti,ab.
11	8 or 9 or 10
12	exp animals/
13	humans.sh.
14	12 not 13
15	7 and 11
16	15 not 14
17	Limit 16 to English language

49	#48 NOT [medline]/lim
48	4 AND 46 AND 47
47	5 OR 6 OR 7 OR 8 OR 9 OR 10
46	23 OR 45
45	35 AND 44
44	36 OR 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43
43	'vs.':ab
42	'vs.':ti
41	'versus':ab
40	'versus':ti
39	'effective*':ab
38	'effective*':ti
37	'compar*':ab
36	'compar*':ti
35	24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34
34	'cross-over study'/de OR 'cross-over study'
33	'matched-pair analysis'/de OR 'matched-pair analysis'
32	'case* and control*':ab
31	'case* and control*':ti
30	'cohort*':ab
29	'cohort*':ti
28	'case control study'/de OR 'case control study'
27	'follow-up'/de OR 'follow-up'
26	'prospective study'/de OR 'prospective study'
25	'longitudinal study'/de OR 'longitudinal study'
24	'cohort analysis'/de OR 'cohort analysis'
23	22 AND 13
22	21 NOT (18 OR 20)
21	11 OR 12
20	19 NOT 14
19	16 OR 17
18	15 NOT 14
17	'random sampl*':ti OR 'random digit*':ti OR 'random effect*':ti OR 'random survey':ti OR 'random regression':ab
16	'random sampl*':ti OR 'random digit*':ti OR 'random effect*':ti OR 'random survey':ti OR 'random regression':ti
15	book:pt OR editorial:pt OR letter:pt OR review:pt
14	'randomized controlled trial'/de OR 'randomized controlled trial'
13	[humans]/lim
12	random*:ti OR placebo*:ti OR 'single blind*':ti OR 'double blind*':ti OR 'triple blind*':ab
11	random*:ti OR placebo*:ti OR 'single blind*':ti OR 'double blind*':ti OR 'triple blind*':ti
10	'atezolizumab':ti OR 'atezolizumab':ab
9	'pembrolizumab':ti OR 'pembrolizumab':ab
8	'nivolumab':ti OR 'nivolumab':ab
7	'afatinib':ti OR 'afatinib':ab
6	'gefitinib':ti OR 'gefitinib':ab
5	'erlotinib':ti OR 'erlotinib':ab
4	1 OR 2 OR 3
3	lung:ab AND (cancer*:ab OR carcin*:ab OR neoplasm*:ab OR tumour*:ab OR tumor*:ab) AND ('non small':ab OR
5	nonsmall:ab) AND cell:ab
2	lung:ti AND (cancer*:ti OR carcin*:ti OR neoplasm*:ti OR tumour*:ti OR tumor*:ti) AND ('non small':ti OR nonsmall:ti)
-	AND cell:ti
1	'non small cell lung carcinoma'/de
-	

Table A4. Search strategy of Embase on June 8, 2016

We performed screening at both the abstract and full-text level. Two investigators screened all abstracts identified through electronic searches according to the inclusion and exclusion criteria described earlier. We did not exclude any study at abstract-level screening due to insufficient information. For example, an abstract that did not report an outcome of interest would be accepted for further review in full text. We retrieved the citations that were accepted during abstract-level screening for full text appraisal. Two investigators reviewed full papers and provided justification for exclusion of each excluded study.

We used criteria modified slightly from those published by the US Preventive Services Task Force (USPSTF) to assess the quality of RCTs and comparative cohort studies, using the categories "good," "fair," or "poor" as described below:^{92,93}

Good: Meets all criteria: Comparable groups were assembled initially and maintained throughout the study; reliable and valid measurement instruments were used and applied equally to the groups; interventions were spelled out clearly; all important outcomes are considered; and appropriate attention paid to confounders in analysis. In addition, for RCTs, at least modified intention to treat (mITT) analysis was done for RCTs.⁹⁴

Fair: Studies were graded "fair" if any or all of the following problems occurred, without the fatal flaws noted in the "poor" category: Generally comparable groups are assembled initially but some question remains whether some (although not major) differences occurred with follow-up; measurement instruments were acceptable (although not the best) and generally applied equally; some but not all important outcomes were considered; and some but not all potential confounders were addressed.

Poor: Studies were graded "poor" if any of the following fatal flaws existed: Groups assembled initially were not close to being comparable or maintained throughout the study; unreliable or invalid measurement instruments were used or not applied equally among groups (including not masking outcome assessment); and key confounders were given little or no attention. For RCTs, intention to treat analysis was lacking.

Appendix B. PRISMA and Evidence Review Table

Figure B1. PRISMA flow Chart Showing Results of Literature Search for NSCLC

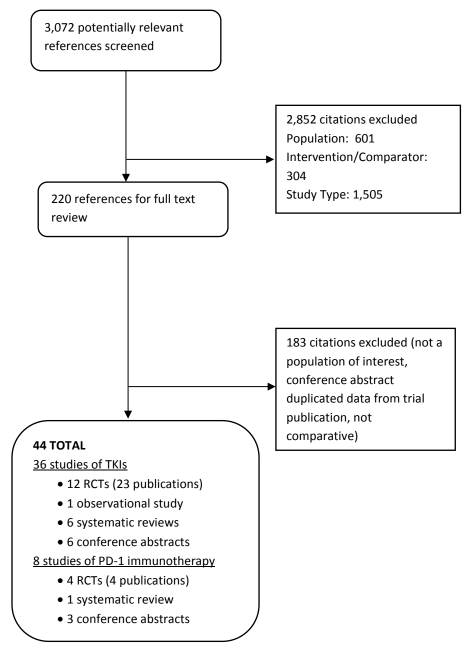


Table B1. Summary evidence table

Author & Year of Publication (Trial) <i>Quality rating</i>	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT	1) Afatinib (n=230)	Treatment-naive	Age, median (range)	Median PFS, m	AEs ≥3, n (%)
	Multicenter		advanced lung	1) 61.5 (28-86)	1) 11.14	Diarrhea
Sequist LV	Open-label	2) Cisplatin +	adenocarcinoma;	2) 61.0 (31-83)	2) 6.90	1) 33 (14.4)
J Clin Oncol 2013	Phase III	pemetrexed (n=115)	harboring EGFR		HR=0.58	2) 0
			mutation;	Male, n (%)	95% CI 0.43-0.78	
(LUX-Lung 3) ³⁵	Primary data cutoff	Dosing schedule	ECOG PS 0-1;	1) 83 (36.1)	p<0.001	Rash/acne
	median follow-up:	1) 40 mg/day	adequate end-organ	2) 38 (33.0)		1) 37 (16.2)
Fair	16.4 m	2) 75 mg/m ² + 500	function;		25 th percentile OS, m	2) 0
		mg/m ² once every 21	measurable disease	White, n (%)	(immature)	
	Location	days up to 6 cycles	(RECIST vs. 1.1)	1) 61 (26.5)	1) 16.6	Stomatitis/mucositis
	133 centers in 25			2) 30 (26.1)	1) 14.8	1) 20 (8.7)
	countries in Asia,				HR=1.12	2) 1 (0.9)
	Europe, North			East Asian, n (%)	95% CI 0.73-1.73	
	America, South			1) 165 (71.7)	p=0.60	Paronychia
	America, and Australia			2) 83 (72.2)		1) 26 (11.4)
					Time to deterioration for	2) 0
				ECOG PS=1, n (%)	worsening of cough	
				1) 138 (60.0)	HR=0.60	Fatigue
				2) 73 (63.5)	95% CI 0.41-0.87	1) 3 (1.3)
					p=0.007	2) 14 (12.6)
				Stage IV, n (%)		
				1) 210 (91.3)	Dyspnea	Treatment-related death
				2) 98 (85.2)	HR=0.68	1) 4
					95% CI 0.50-0.93	2) 0
				Smoker (never), n (%)	p=0.01	
				1) 155 (67.4)		Discontinuation d/t AEs
				2) 81 (70.4)	Pain	1) 23 (10.0)
					NS	2) 17 (15.3)

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	See Sequist LV	See Sequist LV	See Sequist LV	See Sequist LV	QLQ-C30, mean treatment	
	J Clin Oncol 2013	J Clin Oncol 2013	J Clin Oncol 2013	J Clin Oncol 2013	difference (95% CI)	
Yang JC-H						
J Clin Oncol 2013		European Organization			Global health status/QoL	
		for Research and			-3.18 (-5.75 to -0.61)	
(LUX-Lung 3) ⁶¹		Treatment of Cancer			p=0.015	
		Quality of Life				
Fair		Questionnaire C30 and			Physical	
		Lung Cancer-13			-4.80 (-7.47 to -2.13)	
		questionnaires			p<0.001	
					Role	
					-4.40 (-7.40 to -1.40)	
					p=0.004	
					P	
					Emotional NS	
					Cognitive	
					-3.16 (-5.47 to -0.85)	
					p=0.007	
					p=0.007	
					Social NR	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Wu Y-L	RCT Multicenter Open-label	1) Afatinib (n=242) 2) Cisplatin +	EGFR mutation- positive, treatment- naïve, stage IIIB or	Age, median(range) 1) 58 (49-65) 2) 58 (49-62)	Median PFS, m (95% Cl) 1) 11.0 (9.7-13.7) 2) 5.6 (5.1-6.7)	AEs ≥3, n (%) Diarrhea 1) 13 (5.4)
Lancet Oncol 2014	Phase III	gemcitabine (n=122)	IV lung adenocarcinoma;	Male, n (%)	HR=0.28 95% Cl 0.20-0.39	2) 0
(LUX-Lung 6) ³⁷ Fair	Median follow-up (PFS): 16.6 m (IQR 4.7- 9.4)	Dosing schedule 1) 40 mg/day 2) 75 mg/m ² on day 1 +	ECOG PS 0–1; measurable disease (RECIST vs. 1.1);	1) 87 (36.0) 2) 39 (32.0)	p<0.0001 ORR, n (%)	Rash or acne 1) 35 (14.6) 2) 0
	Location 36 centers in China, Thailand, and South Korea	1000 mg/m ² on day 1 and day 8, in a 3-week schedule for maximum of 6 cycles	adequate organ function	Asian, % 1) 100 2) 100 ECOG PS=1, n (%) 1) 194 (80.2) 2) 81 (66.4)	1) 162 (66.9) 2) 28 (23.0) OR=7.28 95% CI 4.36-12.18 p<0.0001 Median OS, m (95% CI)	Stomatitis or mucositis 1) 13 (5.4) 2) 0 Vomiting 1) 2 (0.8)
				Stage IV, n (%) 1) 226 (93.4) 2) 116 (95.1)	(immature) 1) 22.1 (20.0-NE) 2) 22.2 (18.0-NE) HR =0.95 95% CI 0.68-1.33	2) 22 (19.5) Anemia 1) 1 (0.4) 2) 10 (9)
				Smoker (never), n (%) 1) 181 (74.8) 2) 99 (81.1) Uncommon EGFR	p=0.76 Improvement in overall health status/QOL, n (%) 1) 143 (62.7)	Treatment-related death 1) 1 2) 1
				mutation, n (%) 1) 26 (10.7) 2) 14 (11.5)	2) 33 (32.7) p<0.0001	Discontinuation d/t AEs 1) 21 (8.8) 2) 45 (39.8)

Author & Year of Publication (Trial) <i>Quality rating</i>	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Geater SL J Thorac Oncol 2015 (LUX-Lung 6) ⁶³ Fair	See Wu Y-L Lancet Oncol 2014	See Wu Y-L Lancet Oncol 2014 QLQ-C30	See Wu Y-L Lancet Oncol 2014	See Wu Y-L Lancet Oncol 2014	Significantly greater improvements with afatinib in global health status/QoL, physical functioning, role functioning, and social functioning Longer time to deterioration with afatinib vs. cisplatin/gemcitabine for cough, dyspnea, and pain, and all functioning scales and global health status/QoL Improvements in mean scores with afatinib vs. cisplatin/gemcitabine for cough, dyspnea, and pain. Better mean scores over time for all functioning scales, and global health status/QoL	See Wu Y-L Lancet Oncol 2014

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	See Sequist LV J Clin Oncol 2013 and	See Sequist LV J Clin Oncol 2013 and	See Sequist LV J Clin Oncol 2013	See Sequist LV J Clin Oncol 2013 and	Median OS, m <u>LUX-Lung 3</u>	See Sequist LV J Clin Oncol 2013 and
Yang J C-H Lancet	Wu Y-L	Wu Y-L	and	Wu Y-L	1) 28.2 (24.6-33.6)	Wu Y-L
Oncol 2015	Lancet Oncol 2014	Lancet Oncol 2014	Wu Y-L Lancet Oncol 2014	Lancet Oncol 2014	2) 28.2 (20.7-33.2) HR=0.88	Lancet Oncol 2014
(LUX-Lung 3 and LUX-	Data cutoff				95% CI 0.66-1.17	
Lung 6) ³⁶	<u>LUX-Lung 3:</u> Nov 14, 2013				p=0.39	
Fair	<u>LUX-Lung 6:</u> Dec 27, 2013 <i>Median follow-up</i> <u>LUX-</u> <u>Lung 3:</u> 41 m (IQR 35- 44) <u>LUX-Lung6:</u> 33 m (IQR 31-37)				LUX-Lung 6 1) 23.1 (20.4-27.3) 2) 23.5 (18.0-25.6) HR=0.93 95% CI 0.72-1.22 p=0.61	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT	1) Afatinib (n=160)	Age ≥18; common	Age, median (range)	Median PFS, m (95% CI)	AEs ≥3, n (%)
	Multicenter		EGFR mutation;	1) 63 (30-86)	1) 11.0 (10.6-12.9)	Diarrhea
Park K	Open-label	2) Gefitinib	treatment-naïve;	2) 63 (32-89)	2) 10.9 (9.1-11.5)	1) 20 (13)
Lancet Oncol 2016	Phase IIb	(n=159)	stage IIIB or IV lung		HR=0.73	2) 2 (1)
			adenocarcinoma;	Male, n (%)	95% CI 0.57-0.95	
(LUX-Lung 7) ³⁸	Median follow-up	Dosing schedule	ECOG PS 0-1;	1) 69 (43)	p=0.017	Rash/acne
	(PFS): 27.3 m	1) 40 mg/day	measurable disease	2) 53 (33)		1) 15 (9)
Good		2) 250 mg/day	(RECIST vs. 1.1);		Median OS, m (95% CI)	2) 5 (3)
	Location		adequate organ	Asian, n (%)	(immature)	
	64 sites in Australia,		function	1) 94 (59)	1) 27.9 (25.1-32.2)	Fatigue
	Canada, China, France,			2) 88 (55)	2) 25.0 (20.6-29.3)	1) 9 (6)
	Germany, Ireland,				HR=0.87	2) 0
	Norway, Republic of			ECOG PS=1, n (%)	95% CI 0.66-1.15	
	Korea, Singapore,			1) 109 (68)	p=0.33	Increased ALT/AST
	Spain, Sweden,			2) 112 (70)		1) 0
	Taiwan, and United				ORR, n (%)	2) 14 (9)
	Kingdom			Stage IV, n (%)	1) 112 (70)	
	5			1) 152 (95)	2) 89 (56)	Treatment-related death
				2) 156 (98)	OR=1.87	1) 0
				, ()	95% CI 1.18-2.99	2) 1
				Smoker (never), n (%)	p=0.0083	,
				1) 106 (66)	P	Discontinuation d/t AEs
				2) 106 (67)		1) 18 (11)
				_, (0. ,		2) 17 (11)
				EGFR mutation, %		-, -, (,
				L858R 42		
				Del19 58		

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT Multicenter	Overall study population	Age ≥18; stage IIIB or IV	Overall study population Age, median (range)	PFS HR=0.48	Overall study population
Mok TS N Engl J Med 2009	Open-label Phase III Median follow-up	1) Gefitinib (n=609) 2) Carboplatin + paclitaxel (n=608)	lung adenocarcinoma; nonsmokers (<100 cigarettes in	1) 57 (24-84) 2) 57 (25-84) Stage IV, n (%)	95% CI 0.36-0.64 p<0.001 OS	Discontinuation d/t AEs 1) 6.9% 2) 13.6%
(IPASS) ³³	(PFS): 5.6 m	EGFR mutation, n (%)	lifetime) or former light smokers	1) 459 (75) 2) 463 (76)	HR=0.78 95% CI 0.50-1.20	
Fair	Location 87 centers in Hong Kong, elsewhere in China, Indonesia, Japan, Malaysia, the Philippines, Singapore, Taiwan, and Thailand	L858R 42 or Del19 58 1) 130 (98) 2) 121 (99) <u>Dosing schedule</u> 1) 250 mg/day 2) carboplatin at a dose calculated to produce an AUC of 5-6 mg/ml per min over 15-60 min + 200 mg/m ² paclitaxel on day one, every 21 days up to 6 cycles	(stopped smoking ≥15 years previously and had total of ≤10 pack-years of smoking); no previous chemotherapy or biologic or immunologic therapy; ECOG PS 0- 2; measurable disease (RECIST vs. 1.1); adequate hepatic function and neutrophil count	EGFR+ population Male, n (%) 1) 24 (18.2) 2) 26 (20.2) Asian, % 100 ECOG PS 0/1, n (%) 1) 119 (90.2) 2) 122 (94.6) Smoker (never), n (%) 1) 124 (93.9) 2) 122 (94.6)	p=0.33 ORR, n (%) 1) 94 (71.2) 2) 61 (47.3) OR=2.75 95% CI 1.65-4.60 p<0.001 Sustained clinically relevant improvement, % $FACT-L TOI LCS$ 1) 77 70 76 2) 45 38 54 'Trial Outcome Index (sum of physical and functional wellbeing, and lung-cancer subscale)	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Fukuoka M J Clin Oncol 2011 (IPASS) ³⁹ Fair	See Mok TS N Engl J Med 2009 Median follow-up (OS): 17.0 m	See Mok TS N Engl J Med 2009	See Mok TS N Engl J Med 2009	See Mok TS N Engl J Med 2009	Median OS, m 1) 21.6 2) 21.9 HR=1.00 95% CI 0.76-1.33 p=0.99 Median PFS, m 1) 9.5 2) 6.3 HR=0.48 95% CI 0.36-0.64 p<0.001	See Mok TS N Engl J Med 2009
Publication Thongprasert S J Thorac Oncol 2011 (IPASS) ⁹⁵ Fair	See Mok TS N Engl J Med 2009	See Mok TS N Engl J Med 2009	See Mok TS N Engl J Med 2009	See Mok TS N Engl J Med 2009	Median Toxicity-free survival, m 1) 12.1 2) 0.5 HR=0.29 95% CI 0.21-0.39 p<0.001 Time to worsening in HrQoL, m (95% CI) FACT-L 1) 15.6 (11.0-NC) 2) 3.0 (1.5-5.3) TOI 1) 16.6 (11.1-NC) 2) 2.9 (1.5-7.0) LSC 1) 11.3 (11.0-NC) 2) 2.9 (2.1-6.9)	See Mok TS N Engl J Med 2009

Author & Year of Publication (Trial) <i>Quality rating</i>	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Maemondo M	Multicenter Open-label Phase III	1) Gefitinib (n=114) 2) Carboplatin +	Presence of advanced non- small-cell lung	Age, mean (range) 1) 63.9 (43-75) 2) 62.6 (35-75)	Median PFS, m (95% Cl) 1) 10.8 2) 5.4	AEs ≥3, n (%) Neuropathy
N Engl J Med 2010	Median follow-up: 527	paclitaxel (n=114)	cancer harboring sensitive	Male, n (%)	HR=0.30 95% CI 0.22-0.41	1) 0 2) 7 (6.2)
(NEJ002) ⁴¹	days (>17 m)	Dosing schedule 1) 250 mg/day (until	EGFR mutations; absence	1) 42 (36.8) 2) 41 (36.0)	p<0.001	Arthralgia
Fair	Location 43 institutions in Japan	progression, intolerable toxicity, or withdrawal of consent) 2) carboplatin at a dose calculated to produce an AUC of 6 over 1 hr + 200 mg/m ² paclitaxel over 3 hrs, on day 1 every 21 days, for at least 3 cycles	of T790M; no history of chemotherapy; age ≤ 75	Asian, % 100 ECOG PS=1, n (%) 1) 59 (51.8) 2) 55 (48.2) Stage IV, n (%) 1) 88 (77.2) 2) 84 (73.7) Smoker (never), n (%) 1) 75 (65.8) 2) 66 (57.9) L858R: 42.5% Del19: 51.3%	ORR, n (%) 1) 84 (73.7) 2) 35 (30.7) p<0.001 Median OS, m 1) 30.5 2) 23.6 p=0.31 2-year survival, % 1) 61.4 2) 46.7	1) 1 (0.9) 2) 8 (7.1) AST/ALT elevation 1) 30 (26.3) 2) 1 (0.9) Neutropenia 1) 1 (0.9) 2) 74 (65.5) Anemia 1) 0 2) 6 (5.3)

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Oizumi S Oncologist 2012 (NEJ002) ⁶⁴ Fair	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010 Deterioration: score change from baseline by one of 11 points (9.1%) in a direction indicating a worse QoL at any timepoint	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010	Significant differences between treatment arms in deterioration of pain and shortness of breath (HR 0.34; 95% CI, 0.23– 0.50; p<0.0001) and daily functioning (HR 0.43; 95% CI, 0.28 – 0.65; p<0.0001); difference in anxiety between arms NS Gefitinib superior on physical well-being scale (p<0.0001), daily functioning (p=0.035), and subjective QoL (p=0.042).	See Maemondo M N Engl J Med 2010
Publication Inoue A Ann Oncol 2013 (NEJ002) ⁴⁰ Fair	See Maemondo M N Engl J Med 2010 Median follow-up: 704 days	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010	Median OS, m Updated in Dec 2009 1) 27.7 2) 26.6 HR=0.887 95% Cl 0.634-1.241 1-yr survival rate 1) 85.0% 2) 86.8% 2-yr survival rate 1) 57.9% 2) 53.7%	See Maemondo M N Engl J Med 2010

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Watanabe S J Thorac Oncol 2014 (NEJ002) ⁹⁶ Fair	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010	See Maemondo M N Engl J Med 2010 Common EGFR mutation, n 1) 109 2) 106 Uncommon EGFR mutation, n 1) 5 2) 5	Common mutation Median OS, m 1) 29.3 2) 28.0 p=0.378 Median PFS, m 1) 11.4 2) 5.4 ORR, % 1) 76 2) 32 Uncommon mutation Median OS, m 1) 11.9 2) 22.8 p=0.102 Median PFS, m 1) 2.2 2) 5.9 ORR, % 1) 20 2) 20	See Maemondo M N Engl J Med 2010

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Mitsudomi T Lancet Oncol 2010 (WJTOG3405) ⁴² <i>Fair</i>	Multicenter Open-label Phase III Median follow-up: 81 days (74-1253) Location 36 centers in Japan	1) Gefitinib (n=86) 2) Cisplatin + docetaxel (n=86) Dosing schedule 1) 250 mg/day 2) 80 mg/m ² over 90 min + 60 mg/m ² docetaxel over 1 hr, once every 21 days for 3-6 cycles	Advanced or recurrent NSCLC; harboring activating EGFR mutations; age ≤75; ECOG PS 0- 1 measurable or non- measurable disease (RECIST); adequate organ function.	Age, median (range) 1) 64 (34-74) 2) 64 (41-75) Male, n (%) 1) 27 (31) 2) 26 (30) Asian, % 100 ECOG PS=1, n (%) 1) 30 (35) 2) 34 (40) Stage IV, n (%) 1) 41 (48) 2) 41 (48) Smoker (never), n (%) 1) 61 (71) 2) 57 (66) L858R 49 Del19 51	Median PFS, m (95% Cl) 1) 9.2 (8.0-13.0) 2) 6.3 (5.8-7.8) HR=0.489 95% Cl 0.336-0.710 p<0.0001 Median OS, m (95% Cl) 1) 30.9 (24.1-NE) 2) Not reached (15.0-NE) HR=1.638 95% Cl 0.749-3.582 p=0.211 ORR, n (%) 1) 36 (62.1) 2) 19 (32.2) P<0.0001	AEs ≥3, n (%) AST 1) 14 (16) 2) 1 (0.1) ALT 1) 24 (32) 2) 2 (0.2) Leukocytopenia 1) 0 2) 43 (49) Neutropenia 1) 0 2) 74 (19) Anemia 1) 0 2) 15 (17)
Abstract Mitsudomi T J Clin Oncol 2012 (WJTOG3405) ⁹⁷	See Mitsudomi T Lancet Oncol 2010 Median follow-up: 34 m	See Mitsudomi T Lancet Oncol 2010	See Mitsudomi T Lancet Oncol 2010	See Mitsudomi T Lancet Oncol 2010	Median OS, m (95% CI) 1) 36 (26.3 -) 2) 39 (31.2 -) HR=1.185 95% CI 0.767-1.829	See Mitsudomi T Lancet Oncol 2010
Abstract Yoshioka H J Clin Oncol 2014 (WJTOG3405) ⁴³	See Mitsudomi T Lancet Oncol 2010 Median follow-up: 59.1 m	See Mitsudomi T Lancet Oncol 2010	See Mitsudomi T Lancet Oncol 2010	See Mitsudomi T Lancet Oncol 2010	Median OS, m (95% CI) 1) 34.8 (26.0 -39.5) 2) 37.3 (31.2 -45.5) HR=1.252 95% CI 0.883-1.775	See Mitsudomi T Lancet Oncol 2010

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Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Han J-Y J Clin Oncol 2012 (First-SIGNAL) ³² Fair	Multicenter Open-label Phase III Median follow-up: 34 m <u>Location</u> 3 major hospitals in Korea	Overall study population 1) Gefitinib (n=159) 2) Cisplatin + gemcitabine (n=150) EGFR+ 1) Gefitinib (n=26) 2) Cisplatin + gemcitabine (n=16) Dosing schedule 1) 250 mg/day 2) 80 mg/m ² every 21 days + 1250 mg/m ² paclitaxel on day 1 and 8 every 21 up to 9 cycles	Chemotherapy- naïve; never- smoker; age >18 years; stage IIIB or IV lung adenocarcinoma; measurable or nonmeasurable disease; ECOG PS 0-2; adequate bone marrow, liver, and renal function	Overall study population Age, median (range) 1) 57.0 (32-74) 2) 56.5 (19-74) Male, n (%) 1) 19 (12) 2) 16 (11) Asian, % 100 ECOG PS=1, n (%) 1) 104 (65) 2) 105 (70) Stage IV, n (%) 1) 142 (89) 2) 136 (91)	EGFR+ subgroup Median OS, m 1) 27.2 2) 25.6 HR=1.043 95% Cl 0.498-2.182 PFS 1) 8.0 2) 6.3 HR=0.544 95% Cl 0.269-1.100 p=0.086 ORR, n (%) 1) 22 (84.6) 2) 6 (37.5) p=0.002 QLQ-C30 NS LC13 NS	NR for EGFR+ Overall study population AEs ≥3, n (%) Any 1) 46 (28.9) 2) 102 (68) Treatment-related death 1) 0 2) 1
Poster presentation Singh C J ImmunoTher Cancer 2014 ⁹⁸	RCT	 Gefitinib (n=30) Cisplatin + paclitaxel (n=30) <u>Dosing schedule</u> 1) 250 mg/day 2) NR 	Metastatic non- small-cell lung cancer and EGFR mutations who had not previously received chemotherapy	NR	Median PFS, m 1) 10 2) 5 Median OS, m 1) 30 2) 24 ORR, % 1) 70 2) 30	NR

Author & Year of Publication (Trial) <i>Quality rating</i>	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT	1) Erlotinib (n=86)	Stage IIIb or IV	Age, median (range)1) 65	Median OS, m (95% CI)	AEs ≥3, n (%)
	Multicenter		NSCLC; measurable	(24-82)	1) 19.3 (14.7-26.8)	Fatigue
Rosell R	Open-label	2) Cisplatin* +	disease; activating	2) 65 (29-82)	1) 19.5 (16.1-NE)	1) 5 (6)
Lancet Oncol	Phase III	docetaxel or	EGFR mutations;		HR =1.04	2) 16 (20)
2012		gemcitabine (n=87)	age ≥18; no history	Female, n (%)	95% CI 0.65-1.68	p=0.0086
	Median follow-up:		of chemotherapy	1) 58 (67)	p=0.87	
(EURTAC) ⁴⁵	1) 18.9 m	Dosing schedule	for metastatic	2) 68 (78)		Rash
	2) 14.4 m	1) 150 mg/day	disease		Median PFS, m (95% CI)	1) 11 (13)
Fair		2) 3 week cycles of 75		ECOG PS=1, n (%)	1) 9.7 (8.4-12.3)	2) 0
		$mg/m^2 + 75 mg/m^2 on$		1) 47 (55)	2) 5.2 (4.5-5.8)	p=0.0007
		day 1 or 75 mg/m ² on		2) 45 (52)	HR=0.37	
		day 1 + 1250 mg/m ² on			95% CI 0.25-0.54	Neutropenia,
		days 1 and 8		Stage IV, n (%)	p<0.0001	thrombocytopenia
				1) 78 (91)		1) 0, 0
		*Patients ineligible for		2) 82 (94)	Rate of PFS at 1 yr, % (95%	2) 18 (22), 12 (15
		cisplatin: carboplatin (3		, , ,	CI)	p<0.0001, p=0.0003
		wk cycles of AUC 6 on		Never smoked, n (%)	1) 40 (28-52)	
		day 1 with 75 mg/m ²		1) 57 (66)	2) 10 (4-20)	Treatment-related death
		docetaxel on day 1, or		2) 63 (72)		1) 1 (1)
		AUC 5 on day 1 with		, (2) 2 (2)
		1000 mg/m ²		Adenocarcinoma, n (%)		, , ,
		gemcitabine on days 1		1) 82 (95)		Discontinuation d/t AE
		and 8)		2) 78 (90)		1) 11 (13)
				-,		2) 19 (23)
						2) 19 (23)

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Costa C Clin Cancer Res 2014 (EURTAC) ⁴⁴ Fair	See Rosell R Lancet Oncol 2012 Updated results Median follow-up as of January 25, 2013 1) 40.7 m 2) 22.1 m Crossover permitted at time of progression; 80% of chemotherapy group received erlotinib	See Rosell R Lancet Oncol 2012 1) n=50 2) n=45	Subanalysis of 95 patients for whom pretreatment tumor specimens were available	Age <65, n (%) 1) 23 (46) 2) 21 (47) Female, n (%) 1) 34 (68) 2) 37 (82) ECOG PS=1, n (%) 1) 27 (54) 2) 24 (53) Stage IV, n (%) 1) 44 (88) 2) 43 (96) Never smoked, n (%) 1) 32 (64) 2) 32 (71) Adenocarcinoma, n (%) 1) 47 (94) 2) 40 (89) T790M mutation, n (%) 1) 34 (68) 2) 28 (62)	Median overall survival (overall pop. EURTAC), m 1) 22.9 2) 22.1 Median PFS Overall pop. EURTAC, m 1) 10.4 2) 5.1 HR=0.33 95% Cl 0.23-0.49 p<0.0001 No T790M mutation, m (95% Cl) 1) 15.8 (8.8-NR) 2) 5.1 (1.1-6.7) With T790M mutation, m (95% Cl) 1) 9.7 (6.9-12.9) 2) 6.0 (4.1-7.7)	See Rosell R Lancet Oncol 2012

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT	1) Erlotinib (n=110)	Age ≥18 years;	Age, median (range)1)	Median OS, m	AEs ≥3, n (%)
	Multicenter		stage IIIb/IV EGFR+	57.5 (33-79)	1) 26.3	
Wu YL	Open-label	2) Gemcitabine +	NSCLC; EGOG PS 0-	2) 56.0 (30-78)	1) 25.5	Neutropenia
Ann Oncol	Phase III	cisplatin (n=107)	2; no prior exposure		HR=0.91	1) 1 (1)
2015			to chemotherapy or	Male, %	95% CI 0.63-1.31	2) 26 (25)
(510) 05146	Median duration of	Dosing schedule	agents targeting	1) 38.2	p=0.607	
(ENSURE) ⁴⁶	follow-up:	1) 150 mg/day until	HER receptors; no	2) 39.3		Anemia
_ .	1) 28.9 m	progression/	brain metastases		Median PFS, m (95% CI)	1) 1 (1)
Fair	2) 27.1 m	unacceptable toxicity		ECOG PS=1, %	1) 11.0	2) 13 (13)
		2) 1250 mg/m ² days 1		1) 78.9	2) 5.5	
	Location	and 8 + 75 mg/m ² day		2) 79.8	HR=0.34	Leukopenia
	China, Malaysia,	1, every 3 weeks, up to			95% CI 0.22-0.51	1) 1 (1)
	Philippines	4 cycles		Stage IV, %	p<0.0001	2) 15 (14)
				1) 90.9		
	Interim analysis			2) 93.5	ORR, %	Thrombocytopenia
	(cutoff July 20, 2012)			News and the set of	1) 62.7	1) 0
				Never smoker, %	2) 33.6	2) 7 (7)
				1) 71.8	p=NR	Dh
				2) 69.2		Rash
						1) 7 (6)
				Adenocarcinoma, %		2) 1 (1)
				1) 94.5		
				2) 94.4		Discontinuation d/t AE
				Sausmous coll 0/		1) 3 (3)
				Squamous-cell, %		2) 13 (13)
				1) 1.8		
l				2) 1.9		

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Abstract	See Wu YL Ann Oncol	See Wu YL Ann Oncol	See Wu YL Ann Oncol	See Wu YL Ann Oncol	Median PFS, m 1) 11.0	See Wu YL Ann Oncol
Wu YL J Thorac Oncol 2013 (ENSURE) ⁹⁹	2015 Updated analysis (cutoff November 19, 2012)	2015	2015	2015	2) 5.5 HR=0.33 95% CI 0.23-0.47 p<0.0001 ORR, % 1) 68.2 1) 39.3 p<0.0001 Disease control rate, % 1) 91.8 1) 82.2 p=0.0354	2015

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Abstract	See Wu YL	See Wu YL	See Wu YL	See Wu YL	Time to symptomatic	See Wu YL
	Ann Oncol	Ann Oncol	Ann Oncol	Ann Oncol	progression	Ann Oncol
Wu YL J Thorac Oncol	2015	2015	2015	2015	(≥3-point decline in LCS from	2015
2014					baseline), m	
	Data cutoff November				1) 13.8	
(ENSURE) ¹⁰⁰	19, 2012				2) 5.5	
					HR=0.56	
	QoL analysis based on				95% CI 0.36-0.87	
	Functional Assessment				p=0.0076	
	of Cancer Therapy-					
	Lung (FACT-L) and				Time to deterioration	
	Lung Cancer Subscale				TOI, m	
	(LCS)				1) 11.4	
					2) 4.2	
	Trial outcome index				HR=0.51	
	(TOI; ≥6-point decline				95% CI 0.34-0.76	
	in LCS score + physical				p=0.0006	
	and functional scores					
	from baseline)				QoL, m	
					1) 8.2	
	QoL (≥6-point decline				2) 2.8	
	in TOI + social and				HR=0.64	
	emotional scores from				95% CI 0.44-0.93	
	baseline)				p=0.0168	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT Multicenter	1) Erlotinib (n=82)	Age ≥18 years; stage IIIb/IV EGFR+	Age, median (range)1) 57 (31-74)	OS data not mature at primary analysis	AEs ≥3, n (%)
Zhou C Lancet Oncol 2011 (OPTIMAL) ⁴⁷ <i>Fair</i>	Multicenter Open-label Phase III Median duration of follow-up: 15.6 m Primary analysis data cutoff: August 16, 2010 Location 22 centers in China	2) Gemcitabine + carboplatin (n=72) <u>Dosing schedule</u> 1) 150 mg/day until progression/ unacceptable toxicity 2) 1000 mg/m ² days 1 and 8 + AUC 5 day 1, every 3 weeks, up to 4 cycles	stage IIIb/IV EGFR+ NSCLC; measurable disease (RECIST); EGOG PS 0-2; no prior exposure to systemic anticancer therapy Exclusion: uncontrolled brain metastases	(31-74) 2) 59 (36-78) Male, n (%) 1) 34 (41) 2) 29 (40) ECOG PS=2, n (%) 1) 7 (9) 2) 3 (4) Stage IV, n (%) 1) 71 (87) 2) 67 (93) Non-smoker, n (%) 1) 59 (72) 2) 50 (69) Adenocarcinoma, n (%) 1) 72 (88)	primary analysis Deaths, n (%) 1) 16 (20) 1) 12 (17) Median PFS, m (95% Cl) 1) 13.1 (10.58-16.53) 2) 4.6 (4.21-5.42) HR=0.16 95% Cl 0.10-0.26 p<0.0001 ORR, n (%) 1) 68 (83) 2) 26 (36) p<0.0001 Disease control rate, n (%) 1) 79 (96) 2) 59 (82) p=0.0022	Any event 1) 14 (17) 2) 47 (65) Neutropenia 1) 0 2) 30 (42) Thrombocytopenia 1) 0 2) 29 (40) Anemia 1) 0 2) 9 (13) Treatment-related death 1) 0 2) 0
				2) 62 (86)		Discontinuation d/t AE 1) 1 (1) 2) 4 (6)

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Abstract Zhou C J Thorac Oncol 2011 (OPTIMAL) ¹⁰¹	See Zhou C Lancet Oncol 2011	See Zhou C Lancet Oncol 2011	See Zhou C Lancet Oncol 2011	See Zhou C Lancet Oncol 2011	Clinically-relevant improvement in FACT-L score, % 1) 73 2) 29.6 OR=6.9 95% Cl 3.07-15.48 p<0.0001 Clinically-relevant improvement in LCS score, % 1) 75.7 2) 31.5 OR=6.77 95% Cl 3.04-15.05 p<0.0001	See Zhou C Lancet Oncol 2011

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Chen G Ann Oncol 2013 (OPTIMAL) ¹⁰² Fair	See Zhou C Lancet Oncol 2011 Updated data cutoff January 7, 2011	See Zhou C Lancet Oncol 2011	See Zhou C Lancet Oncol 2011	Mean baseline score FACT-L 1) 94.43 2) 92.89 TOI 1) 55.76 2) 55.52 LCS 1) 18.46 2) 19.52	Updated median PFS, m (95% Cl) 1) 13.7 (10.58-15.28) 2) 4.6 (4.21-5.42 HR=0.164 95% Cl 0.105-0.256 p<0.0001	See Zhou C Lancet Oncol 2011
					p 0.0067 0.003 0.001	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	See Zhou C	See Zhou C	See Zhou C	See Zhou C	Median OS, m	See Zhou C
	Lancet Oncol	Lancet Oncol	Lancet Oncol	Lancet Oncol	1) 22.8	Lancet Oncol
Zhou C	2011	2011	2011	2011	2) 27.2	2011
Ann Oncol					HR=1.19	
2015					95% CI 0.83-1.71	
	Median follow-up for				p=0.2663	
(OPTIMAL) ⁴⁸	OS: 25.9 m					
					Clinical characteristics did	
Fair	Data cutoff: December				not have significant impact	
	21, 2012				on OS	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Gridelli C J Clin Oncol 2012 (TORCH) ³⁴ Fair	RCT Multicenter Open-label Phase III Median duration of follow-up: 24.3 m Location Italy and Canada	 Erlotinib (n=380) Cisplatin + gemcitabine (n=380) <u>Dosing schedule</u> 150 mg/day until progression 80 mg/m² day 1 + 1200 mg/m² on days 1 and 8, every 3 weeks, up to 6 cycles Note: After progression, patients crossed over to opposite treatment arm 	Stage IIIb/IV NSCLC; ≥1 target/nontarget lesion (RECIST); age <70 (no age restrictions for Canadian centers); ECOG PS 0-1; no prior treatment with anti-EGFR agents; adequate bone marrow, hepatic, and renal function; asymptomatic brain metastases eligible	Age, median (range)1) 63 (27-79) 2) 62 (34-81) Male, n (%) 1) 252 (66) 2) 252 (66) ECOG PS=1, n (%) 1) 183 (48) 2) 195 (51) Stage IV, n (%) 1) 334 (88) 2) 343 (90) Never smoker, n (%) 1) 78 (21) 2) 79 21) Adenocarcinoma, n (%) 1) 210 (55) 2) 212 (56) EGFR+, n (%) 1) 19 (14) 2) 20 (15)	Subset of patients with EGFR Mutations Median OS, m (95% CI) 1) 18.1 (12.4-NE) 1) 32.5 (17.3-NE) HR=1.58 95% CI 0.70-3.57 Median PFS, m (95% CI) 1) 9.7 (5.7-18.2) 2) 6.9 (6.6-9.6) HR=0.60 95% CI 0.30-1.20 ORR, % 1) 42.1 2) 25.0	Overall study population AEs \geq 3, n (%) Anemia 1) 17 (5) 2) 30 (8) Neutropenia 1) 1 (0.3) 2) 79 (22) Thrombocytopenia 1) 0 2) 42 (11) Skin rash 1) 40 (11) 2) 1 (0.3)

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	See Gridelli C	See Gridelli C	See Gridelli C	See Gridelli C	EGFR+ patients	See Gridelli C
	J Clin Oncol	J Clin Oncol	J Clin Oncol	J Clin Oncol	Best QoL response	J Clin Oncol
Di Maio M	2012	2012	2012	2012	Global QoL, n (%)	2012
J Thorac Oncol	See Gridelli C	See Gridelli C	See Gridelli C	See Gridelli C	Improved Stable Worse	See Gridelli C
2012	J Clin Oncol	J Clin Oncol	J Clin Oncol	J Clin Oncol	1) 6 (40) 5 4	J Clin Oncol
	2012	2012	2012	2012	(33) (27)	2012
(TORCH) ¹⁰³					2) 8 (50) 4 4 (25) (25)	
	EORTC-C30 and EORTC				Physical Functioning, n (%)	
Fair	QLQ-LC13				Improved Stable Worse	
					1) 5 (33) 6 (40) 4	
	Best QoL response				(27)	
	criteria				2) 8 (50) 3 (19) 5	
	Improved: ≥10 pt				Pain, n (%)	
	improvement since				PdIII, II (70) Improved Stable Worse	
	baseline					
					1) 7 (47) 5 (33) 3 (20)	
	Stable: <10 pt change				2) 9 (56) 4 (25) 3	
	since baseline				(19)	
					Dyspnea, n (%)	
	Worse: ≥10 pt				Improved Stable Worse	
	worsening since				1) 6 (40) 6 3 (40) (20)	
	baseline				(40) (20) 2) 6 (38) 7 3	
					(44) (19)	
					Cough, n (%)	
					Improved Stable Worse	
					1) 7 (47) 5 3	
					(33) (20)	
					2) 7 (44) 5 4 (31) (25)	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Lim SH J Thorac Oncol 2014 ⁴⁹ Fair	Matched-pair case- control Consecutive selection of patients Location Samsung Medical Center, Seoul, Korea	 Gefitinib (n=121) Erlotinib (n=121) Dosing schedule 250 mg/day until progressive disease or unacceptable toxicity 150 mg/day until progressive disease or unacceptable toxicity 	Clinically proven recurrent or advanced/metastati c stage IIIb/IV NSCLC with EGFR mutation; brain metastasis included if underwent whole- brain radiotherapy or stereotactic radiosurgery; first- line or second-line or higher after failure of prior cytotoxic chemotherapy	Age, median (range)1) 58 (29-85) 2) 58 (30-84) Male, n (%) 1) 53 (43.8) 2) 53 (43.8) ECOG PS=0-1, n (%) 1) 110 (91) 2) 110 (91) Stage IV, n (%) 1) 90 (74) 2) 88 (73) Never smoker, n (%) 1) 77 (64) 2) 77 (64) 1 prior regimen, n (%) 1) 65 (54) 2) 82 (68) Adenocarcinoma, n (%) 1) 119 (98) 2) 117 (97)	Subset of patients treated with first-line EGFR TKIs Median OS, m (95% Cl) 1) 24.5 (8.6-40.4) 2) Not reached Median PFS, m (95% Cl) 1) 11.7 (6.7-16.7) 2) 14.5 (8.7-20.4) p=0.507 ORR, % 1) 76.7 2) 90.0 p=0.431	Dose adjustment, n 1) 1 2) 22

Author & Year of Publication (Trial) <i>Quality rating</i>	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Quality rating Publication Fehrenbacher L Lancet 2016 (POPLAR) ²¹ Good	RCTMulticenterOpen-labelPhase IIMedian duration of follow-up 1) 14.8 2) 15.7Primary analysis cutoff date: May 8, 2015Location 13 countries in Europe and N. America	1) Atezolizumab (n=144) 2) Docetaxel (n=143) <u>Dosing schedule</u> 1) 1200 mg until progression or unacceptable toxicity 2) 75 mg/m ² every 3 weeks on day 1 until progression or unacceptable toxicity No crossover permitted	Age ≥18; ECOG PS 0-1; measurable disease (RECIST); adequate hematological and end-organ function; provided tumor specimens for PD-L1 testing Exclusion: Active/untreated CNS metastases; history of pneumonitis, autoimmune or chronic viral diseases	Age, median (range)1) 62 (42-82) 2) 62 (36-84) Male, n (%) 1) 93 (65) 2) 76 (53) ECOG PS=1, n (%) 1) 96 (68) 2) 97 (68) Squamous, n (%) 1) 49 (34) 2) 48 (34) Never smoker, n (%) 1) 27 (19) 2) 29 (20) 1 prior regimen, n (%) 1) 93 (65) 2) 96 (67) EGFR+, n (%) 1) 10 (12) 2) 8 (10)	Median OS, m (95% Cl) 1) 12.6 (9.7-16.4) 2) 9.7 (8.6-12.0) HR=0.73 95% Cl 0.53-0.99 p=0.04 Median PFS, m (95% Cl) 1) 2.7 (2.0-4.1) 2) 3.0 (2.8-4.1) HR=0.94 95% Cl 0.72-1.23 ORR, n (%) 1) 21 (15) 2) 21 (15 Duration of response, m (95% Cl) 1) 14.3 (11.6-NE) 2) 7.2 (5.6-12.5)	AEs ≥3, n (%) Any event 1) 57 (40) 2) 71 (53) Discontinuation d/t AE 1) 11 (8) 2) 30 (22)

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication	RCT Multicenter	1) Nivolumab (n=292)	Documented stage IIIB or	Age, median (range)1) 61 (37-84)	Median OS, m (95% Cl) 1) 12.2 (9.7-15.0)	AEs ≥3, n (%)
Borghaei H N Engl J Med	Open-label Phase III	2) Docetaxel (n=290)	IV or recurrent nonsquamous	2) 64 (21-85)	1) 9.4 (8.1-10.7)	Any event 1) 30 (10)
2015	Interim analysis	Dosing schedule 1) 3 mg/kg every 2	NSCLC after radiation	Male, n (%) 1) 151 (52)	1-yr OS rate, % (95% CI) 1) 51 (45-56)	2) 144 (54)
(CheckMate 057) ⁶⁸	minimum follow-up: 13.2 m	weeks 2) 75 mg/m² every 3	therapy or surgical resection;	2) 168 (58)	2) 39 (33-45) HR=0.73	Neutropenia 1) 0
Good	Additional follow-up minimum: 17.2 m	weeks	disease recurrence or progression during/after one prior platinum-	White, n (%) 1) 267 (91) 2) 266 (92)	96% Cl 0.59-0.89 p=0.002 Median PFS, m (95% Cl)	2) 73 (27) Febrile neutropenia 1) 0
			based doublet chemotherapy	ECOG PS=1, n (%) 1) 208 (71) 2) 194 (67)	1) 2.3 (2.2-3.3) 2) 4.2 (3.5-4.9)	2) 26 (10)
			regimen		Rate of PFS at 1 yr, % (95%	Leukopenia 1) 0 2) 22 (8)
			Primary endpoint: OS	Stage IV, n (%) 1) 272 (93) 2) 266 (92)	CI) 1) 19 (14-23) 2) 8 (5-12) HR=0.92	2) 22 (8) Treatment-related death 1) 1
				Smoker, n (%) 1) 231 (79) 2) 227 (78)	95% Cl 0.77-1.1 p=0.39	2) 1 Discontinuation d/t TEAE
				EGFR+, n (%) 1) 44 (15) 2) 38 (13)	ORR, % (95% Cl) 1) 19 (15-24) 2) 12 (9-17) p=0.02	1) 5% 2) 15%
Abstract	See Borghaei H N Engl J Med	See Borghaei H N Engl J Med	See Borghaei H N Engl J Med	See Borghaei H N Engl J Med	18 m OS rate, % 1) 39	See Borghaei H N Engl J Med
Horn L Eur J Cancer 2015 (CheckMate 057) ⁷²	2015	2015	2015	2015	2) 23 Symptom improvement rate by wk 12, n (%) 1) 52 (18)	2015
					2) 57 (20)	

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Publication Brahmer J N Engl J Med	RCT Multicenter Open-label Phase III	1) Nivolumab (n=135) 2) Docetaxel (n=137)	Stage IIIB or IV squamous-cell NSCLC; disease recurrence after 1	Age, median (range)1) 62 (39-85) 2) 64 (42-84)	Median OS, m (95% Cl) 1) 9.2 (7.3-13.3) 1) 6.0 (5.1-7.3) HR =0.59	AEs ≥3, n (%) Any event 1) 9 (7)
2015	Minimum follow-up:	Dosing schedule 1) 3 mg/kg every 2	prior platinum-containing	Male, n (%) 1) 111 (82)	95% CI 0.44-0.78 p<0.001	2) 71 (55)
(CheckMate 017) ⁶⁹ Good	11 m	weeks 2) 75 mg/m ² every 3 weeks	regimen; age ≥18; ECOG PS <2; submitted	2) 97 (71) White, n (%)	1-yr OS rate, % (95% CI) 1) 42 (34-50)	Fatigue 1) 1 (1) 2) 10 (8)
Good		Weeks	submitted pretreatment tumor-tissue specimen for biomarker analyses Excluded: prior T- cell costimulation, checkpoint-targeted agents, or docetaxel; >1 prior systemic therapy for metastatic disease	White, n (%) 1) 122 (90) 2) 130 (95) ECOG PS=1, n (%) 1) 106 (79) 2) 100 (73) Stage IV, n (%) 1) 105 (78) 2) 112 (82) Smoker, n (%) 1) 121 (90) 2) 129 (94) Squamous, n (%) 272 (100)	1) 42 (34-50) 2) 24 (17-31) Median PFS, m (95% Cl) 1) 3.5 (2.1-4.9) 2) 2.8 (2.1-3.5) HR=0.62 95% Cl 0.47-0.81 p<0.001 1-yr PFS rate, % (95% Cl) 1) 21 (14-28) 2) 6 (3-12) ORR, % (95% Cl) 1) 20 (14-28) 2) 9 (5-15) OR=2.6 (1.3-5.5) p=0.008	2) 10 (8) Neutropenia 1) 0 2) 38 (30) Febrile neutropenia 1) 0 2) 13 (10) Treatment-related death, n 1) 0 2) 3 Discontinuation d/t TEAE, % 1) 3 2) 10

Author & Year of Publication (Trial) <i>Quality rating</i>	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
Abstract Gralla RJ J Thorac Oncol 2015 (CheckMate 017) ⁷³	See Brahmer J N Engl J Med 2015	See Brahmer J N Engl J Med 2015	See Brahmer J N Engl J Med 2015	See Brahmer J N Engl J Med 2015 Baseline LCSS ASBI 1) 29.6 (SD 16.4) 2) 29.6 (SD 14.7)	Week 12 meaningful symptom improvement (≥10 pt decrease), % (95% Cl) 1) 20.0 (13.6-27.7) 2) 21.9 (15.3-29.8) Statistically significant improvements in LCSS ASBI at each assessment (vs. baseline) wks. 12-54 with nivolumab No statistically significant improvement in LCSS ASBI from baseline – wk 18 with docetaxel	See Brahmer J N Engl J Med 2015
Abstract Reck M Ann Oncol 2015 (CheckMate 017) ⁷¹	See Brahmer J N Engl J Med 2015	See Brahmer J N Engl J Med 2015 Minimally important difference (MID) EQ-5D: 0.08 EQ-VAS: 7	See Brahmer J N Engl J Med 2015	See Brahmer J N Engl J Med 2015 Baseline mean EQ-VAS (SD) 1) 63.7 (18.2) 2) 66.3 (20.5) Baseline mean EQ-5D (SD) 1) 0.683 (0.208) 2) 0.663 (0.284)	Nivolumab EQ-VAS at wks. 12, 20 to 36, and 48 higher vs. B/L (p≤0.05); wks. 24 -36 and 48 >MID EQ-5D index at wks. 16 to 30 and wks. 42 to 54 improved vs. B/L (p≤0.05); wks. 42-54 improved vs. B/L (p≤0.05); wks. 42-54 >MID Docetaxel EQ-VAS and EQ-5D index scores did not differ significantly from B/L to wk 18, after which <10 patients in sample	See Brahmer J N Engl J Med 2015

Author & Year of Publication (Trial) Quality rating	Study Design and Duration of F/u	Interventions (n) Dosing schedule	Major Inclusion and Exclusion Criteria	Patient Characteristics	Outcomes	Harms
(Trial)	RCT Multicenter Open-label Phase III Minimum follow-up: 8 m Median Follow-up: 13.1 m			Age, median (range) 1) 63 (56-69) 2) 63 (56-69) 3) 62 (56-69) Male, n (%) 1) 212 (62) 2) 213 (62) 3) 209 (61) White, n (%) 1) 246 (72) 2) 250 (72) 3) 251 (73) ECOG PS=1, n (%) 1) 229 (67) 2) 225 (65) 3) 224 (65) Smoker, n (%) Former or current 1) 279 (81) 2) 285 (82) 3) 269 (78) EGFR+, n (%) 1) 28 (8) 2) 32 (9) 3) 26 (8)	Docetaxel as ref. Median OS, m (95% Cl) 1) 10.4 (9.4-11.9) HR=0.71, 95% Cl 0.58-0.88; p=0.0008 2) 12.7 (10.0-17.3) HR=0.61, 95% Cl 0.49-0.75; p<0.0001 3) 8.5 (7.5-9.8) Median PFS, m (95% Cl) 1) 3.9 2) 4.0 HR=0.88, 95% Cl 0.74-1.05; p=0.07 3) 4.0 <u>PD-L1 TPS\geq50%</u> Median OS, m (95% Cl) 1) 14.9 HR=0.54, 95% Cl 0.38-0.77; p=0.0002 2) 17.3 HR=0.50, 95% Cl 0.36-0.70; p<0.0001 3) 8.2 Median PFS, m (95% Cl) 1) 5.0 HR=0.59, 95% Cl 0.44-0.78; p=0.001	Grade 3-5 AE, n (%) Any 1) 43 (13) 2) 55 (16) 3) 109 (35) Neutropenia 1) 0(0) 2) 0(0) 3) 38 (12) Treatment-related death, n 1) 3 2) 3 3) 5 Discontinuation d/t TEAE, n (%) 1) 15 (4) 2) 17 (5) 3) 31 (10)
					2) 5.2 HR=0.59, 95% CI 0.45-0.78; p<0.001 3) 4.1	

<u>Appendix C. Additional Results from Evidence</u> <u>Review</u>

Table C1. Results of literature search

	Population 1 (1 st -line EGFR+)	Population 2 (1 st -line EGFR-)	Population 3 (2 nd -line EGFR-)	Population 4 (2 nd - or 3 rd -line EGFR+)
Good quality studies, n	1	0	4	0
Fair quality studies, n	23	0	0	0
Poor quality studies, n	0	0	0	0
Not rated, n [*]	12	0	4	0
Total number of relevant references	36	0	8	0

Table C2. EGFR mutations in key TKI trials

Key Trials	1) Treatment	Exon 19 deletion (%)	Exon 21 L858R (%)	Other (%)	
	2) Comparator				
LUX-Lung 3	1) Afatinib	1) 49	1) 40	1) 11	
LOA-Lung 5	2) Cisplatin+Pemetrexed	2) 50	2) 41	2) 10	
LUX-Lung 6	1) Afatinib	1) 51	1) 38	1) 11	
LOX-Lung 0	2) Cisplatin+Gemcitabine	2) 51	2) 38	2) 12	
LUX-Lung 7	1) Afatinib	1) 58	1) 42	NR	
	2) Gefitinib	2) 58	2) 42	NIX	
IPASS∝	1) Gefitinib	1) 50	1) 48	1) 6	
IF ASS	2) Carboplatin+Paclitaxel	2) 57	2) 36	2) 10	
NEJ002	1) Gefitinib	1) 51	1) 43	1) 6	
NLJU02	2) Carboplatin+Paclitaxel	2) 52	2) 42	2) 6	
WJTOG3405	1) Gefitinib	1) 58	1) 42	NR	
1003403	2) Cisplatin+Docetaxel	2) 43 2) 57			
FIRST SIGNAL	1) Gefitinib	64 ^β	36 ^β	NR	
	2) Cisplatin+Gemcitabine		50		
	1) Erlotinib	1) 66	1) 34		
EURTAC	2) Cisplatin+Gemcitabine/	2) 67	2) 33	NR	
	Docetaxel				
ENSURE	1) Erlotinib	1) 52	1) 48	NR	
	2) Cisplatin+Gemcitabine	2) 57	2) 43		
OPTIMAL	1) Erlotinib	1) 52	1) 48	NR	
	2) Carboplatin+Gemcitabine	2) 54	2) 46		
TORCH	1) Erlotinib	NR	NR	NR	
	2) Cisplatin+Gemcitabine				

 α Eleven patients had multiple mutations and are counted once for each type of mutation they had; β total across treatment arms

Trial	Assigned treatment	Duration of treatment	Median (range)	Crossover after progression (% total)
IPASS	gefitinib	daily until progression	5.6 (0.1 to 22.8)	68.2% Platinum doublets 54.5% carboplatin + paclitaxel
	carboplatin +paclitaxel	every 3 weeks until progression or up to 6 cycles	4.1 (0.7 to 5.8)	64.3% TKI 47.3% gefitinib
	gefitinib	daily until progression	163 days (11 to 885)	65% platinum doublets
First-SIGNAL			6 cycles (1 to 9)	75% TKI
	gefitinib	daily until progression	165 days (22 to 1100)	19.8% platinum doublets
WJTOG3405	carboplatin +paclitaxel	every 3 weeks until progression or 3-6 cycles	64 days (1 to 106) 4 cycles (1 to 6)	59.3% TKI
	gefitinib	daily until progression	308 days (14 to 1219)	45.6% carboplatin- paclitaxel
NEJ002	carboplatin +paclitaxel	every 3 weeks until progression or at least 3 cycles	4 cycles (1 to 7)	93.0% gefitinib
	erlotinib	daily until progression	NR	NR
TORCH	cisplatin +gemcitabine	every 3 weeks until progression or up to 6 cycles	5	NR
OPTIMAL	erlotinib	daily until progression	55.5 weeks (3.1 to 93.0)	35.4% chemotherapy only 59.8% chemotherapy alone or in combination
OF THIMAL	cisplatin +gemcitabine	every 3 weeks until progression or up to 4 cycles	10.4 weeks (1.0 to 18.9) 4 cycles (1 to 6)	36.1 TKI only 69.5% TKI alone or in combination
ENSURE	erlotinib	daily until progression	NR	59.1% at least 1 platinum compounds 54.5% at least 1 antimetabolites
	cisplatin +gemcitabine	every 3 weeks until progression or up to 4 cycles	NR	85.6% TKI 51.9% erlotinib
	afatinib	daily until progression	11.0 months	71% subsequent chemotherapy
LUX LUNG 3	cisplatin +pemetrexed	every 3 weeks until progression or up to 6 cycles	6 cycles (1 to 9)	75% TKI
	afatinib	daily until progression	398 days (IQR 173 to 537)	41.7% cisplatin + gemcitabine
LUX LUNG 6	cisplatin +gemcitabine	every 3 weeks until progression or up to 6 cycles	89 days (IQR 60 to 119) 4 cycles	48.4% TKI
	erlotinib	daily until progression	8.2 months (0.3 to 32.9)	37.2% chemotherapy
EURTAC	PD	every 3 weeks until progression or up to 4 cycles	2.8 months (0.7 to 5.1) 4 cycles (1 to 6)	75.86% TKIs 74.71% erlotinib
	afatinib	daily until progression	13.7 months (IQR 7.4 to 24.3)	37.5% TKI
LUX LUNG 7	gefitinib	daily until progression	11.5 months (IQR 6.2 to 18.8)	49.1% TKI

Additional Quality of Life Results: TKIs

For afatinib, both LUX-Lung 3 and LUX-Lung 6 used the Global Health Status/ Quality of Life and functioning scale domains in the European Organization for Research and Treatment of Cancer (EORTC) Quality of Life Questionnaire C30 (QLQ-C30) questionnaires and measured changes in score, proportion of patients with improvement (≥ 10 points), and time-to-deterioration (time to first instance of worsening \geq 10 points). Each scale was standardized to a range from 0 to 100 and a 10-point change was used as the threshold for clinical importance.¹⁰⁵ The longitudinal analysis of LUX-Lung 3 showed that compared with patients on cisplatin plus pemetrexed, patients on afatinib had significantly better scores over time on Global Health Status/QoL (p=0.015), physical (p<0.001), role (p=0.004), and cognitive (p=0.007) functioning. However, measurements of the clinical importance of QoL changes, including proportions of patients with clinically meaningful improvement and time-to-deterioration in QoL and functioning were not significantly different between study arms.⁶¹ However, in LUX-Lung 6, patients on afatinib showed clinically meaningful improvements on the Global Health Status/QoL scale. Specifically, the afatinib arm had a higher proportion of patients with improvement (63% vs. 33%; p<0.0001), and a longer time to deterioration (HR 0.56, 95% CI 0.41 to 0.77; p=0.0002) compared to patients on cisplatin plus gemcitabine. In addition, function scales, including physical, role, emotional, cognitive, and social, were all improved in the afatinib arm compared to the chemotherapy arm.⁶³

Two trials used different instruments to measure the effects of gefitinib on QoL compared to carboplatin plus paclitaxel. The IPASS trial measured quality of life using percentage of patients with sustained clinically relevant improvement on the Functional Assessment of Cancer Therapy-Lung (FACT-L; 0 to 136 scale) and Trial Outcome Index (TOI; 0 to 84 scale) (increase of 6 points or more, maintained for at least 21 days); the TOI is the sum of physical well-being, functional well-being, and lung-cancer subscales on the FACT-L. In EGFR+ patients, gefitinib showed greater proportions of patients improving on both measurements (FACT-L: 70.2% improved vs. 44.5% improved; OR 3.01, 95% CI 1.79 to 5.07; p<0.0001; TOI: 70.2% improved vs. 38.3% improved; OR 3.96, 95% CI 2.33 to 6.71; p<0.0001).³³ Similarly, time-to-worsening, with worsening defined as a decrease of 6 points maintained for at least 21 days, was also longer with gefitinib than with carboplatin/paclitaxel on both FACT-L (15.6 months vs. 3.0 months) and TOI (16.6 months vs. 2.9 months).⁹⁵ The NEJ002 trial assessed benefits of gefitinib compared to carboplatin/paclitaxel on QoL using The Care Notebook questionnaires. The questionnaires contain three major scales, including physical well-being, mental well-being, and life well-being. Patients who received gefitinib were more likely to improve and less likely to get worse (by at least 1 point on a 0-10 scale) compared with patients who received chemotherapy on physical well-being (25% improved and 36% worse with gefitinib vs. 21% improved and 66% worse with chemo; p<0.0001) and life well-being (53% improved and 17% worse with gefitinib vs. 42% improved and 47% worse with chemo; p<0.0001), while no difference was found for mental well-being.64

Two trials compared erlotinib to doublets containing gemcitabine using the same instruments and criteria for clinical importance as used in the IPASS trial. In the ENSURE trial, time-to-deterioration was significantly longer with erlotinib compared to cisplatin-gemcitabine (TOI: 11.4 vs. 4.2 months; HR 0.51, 95% CI 0.34 to 0.76; p=0.0006; FACT-L: 8.2 vs. 2.8 months; HR 0.64, 95% CI 0.44 to 0.93; p=0.0168).¹⁰⁰ In the final analysis of OPTIMAL, a higher proportion of patients on erlotinib achieved improvement in QoL compared to patients on chemotherapy (FACT-L: 74.3% vs. 31.5%; TOI: 73.0% vs. 25.9%).¹⁰²

Table C4. C	Overall survival	by histological	diagnosis
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Drug	Trial	Overall HR	Squamous HR	Non-Squamous HR
Nivolumab	CheckMate 017		0.59 (0.44-0.79)	
Nivolumab	CheckMate 057			0.73 (0.60-0.88)
Atezolizumab	POPLAR	0.73 (0.53-0.99)	0.80 (0.49-1.30)	0.69 (0.47-1.01)
Pembrolizumab	KEYNOTE-010	0.71	0.74 (0.50-1.09)	0.63 (0.50-0.79)

Table C5. PD-L1 tests and cutpoints in the key trials

Trial name	PD-L1 test	Measurement and cutpoints	Subgroups
CheckMate 017	Immunohistochemical assay (Dako North America) using a rabbit antihuman PD-L1 antibody (clone 28-8, Epitomics)	percentage of PD-L1 - expressing tumor cells: ≥1%, ≥5%, and ≥10%	a) $<1\%$ b) $\ge 1\%$ c) $<5\%$ d) $\ge 5\%$ e) $<10\%$ f) $\ge 10\%$
CheckMate 057	same as above	same as above	same as above
KEYNOTE-010	Immunohistochemical assay (Dako North America) using a murine 22C3 antihuman PD-L1 antibody (Merck & Co., Inc.)	percentage of PD-L1 - expressing tumor cells: ≥1% and ≥50%	a) ≥1% b) ≥50%
POPLAR	VENTANA SP142 PD-L1 immunohistochemistry assay (Ventana Medical Systems)	 a) percentage of PD-L1 - expressing tumor cells: TC3≥50%, TC2 ≥5% and <50%, TC1≥1% and <5%, TC0<1% b) percentage of tumor area occupied by PD-L1-expressing tumor-infiltrating immune cells: IC3≥10%, IC2 ≥5% and <10%, IC1≥1% and <5%, IC0<1% 	a) TC3 or IC3 b) TC2/3 or IC2/3 c) TC1/2/3 or IC1/2/3 d) TC0 and IC0

Appendix D. Network Meta-Analysis Methods and Results

Network Meta-Analysis Methods

In addition to summary evidence tables, we performed quantitative indirect comparisons using Bayesian network meta-analysis (NMA) where possible.¹⁰⁶ Results are summarized in the report text. Review of the deviance information criterion (DIC) statistics as well as comparison of the residual deviance (resdev) to the number of unconstrained data points was used to assess the best model fit under multiple alternative assumptions. Given the expectation of at least some degree of heterogeneity in patient populations and/or study design, there is a general preference for a random-effects approach. So we took this approach for TKIs. However, the available network of immunotherapies is constructed of primarily single-study connections, which made the only feasible approach a fixed-effects model (to preserve statistically-significant effects observed in trials) and selected subgroup analyses (to address heterogeneity).¹⁰⁷

Quantitative analyses focused attention on the effects of the regimens of interest on progressionfree and/or overall survival, and were conducted using the NetMetaXL tool (http://www.netmetaxl.com/), a publicly-available and validated Excel-based tool for specifying and analyzing Bayesian indirect comparisons in a WinBUGS environment. For these outcomes, adjusted hazard ratios from the randomized trials were log-transformed and entered into the spreadsheet, and 95% confidence intervals were used to specify variance estimates (i.e., standard errors). A total of 40,000 iterations each were employed for both "burn-in" (for model convergence) and model (for model results) simulations.

We also conducted sensitivity analyses in which digitized information from the progression-free survival and overall survival curves for each immunotherapy trial were used to inform assessment of hazard ratios at multiple timepoints to determine whether the assumption of a proportional hazard holds true, based on established methods.¹⁰⁸. Both Weibull and Gompartz distribution were modeled and whichever better fitted the data was used as input to the economic model. In this instance, 30,000 iterations were used for both burn-in and model simulations.



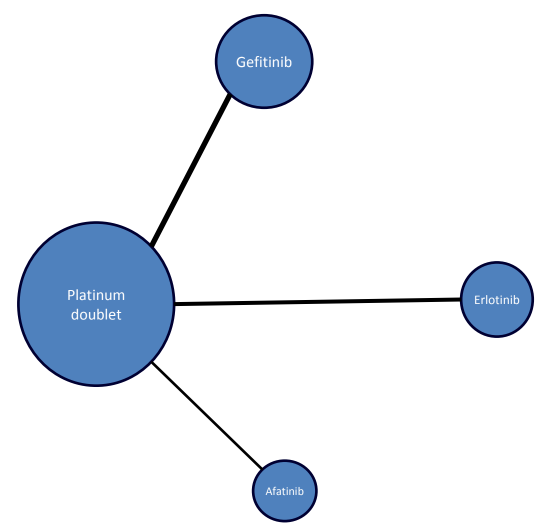


Table D1. Overall survival studies of TKIs (all platinum doublets combined)

Study name	Treatment 1	Treatment 2	n1	n2	LogHR	LogSE
LUX LUNG 3	afatinib	platinum doublets	230	115	-0.12783	0.146051
LUX LUNG 6	afatinib	platinum doublets	242	122	-0.07257	0.134529
IPASS	gefitinib	platinum doublets	132	129	0	0.142759
NEJ002	gefitinib	platinum doublets	114	114	-0.11991	0.171333
WJTOG3405	gefitinib	platinum doublets	86	86	0.224742	0.17812
FIRST-SIGNAL	gefitinib	platinum doublets	26	16	0.042101	0.376887
ENSURE	erlotinib	platinum doublets	110	107	-0.09	0.19
OPTIMAL	erlotinib	platinum doublets	82	72	0.17	0.18
TORCH	erlotinib	platinum doublets	19	20	0.46	0.42

afatinib			RE Model: resdev, 7.029 vs. 9 DIC = -0.829	;
0.91 (0.67 to 1.23)	Platinum doublet		DIG0.029	
0.88 (0.59 to 1.30)	0.97 (0.76 to 1.24)	gefitinib		
0.83 (0.52 to 1.26)	0.91 (0.66 to 1.24)	0.94 (0.61 to 1.40)	erlotinib	

Table D2. Network meta-analysis of TKIs: Overall survival (all platinum doublets combined)



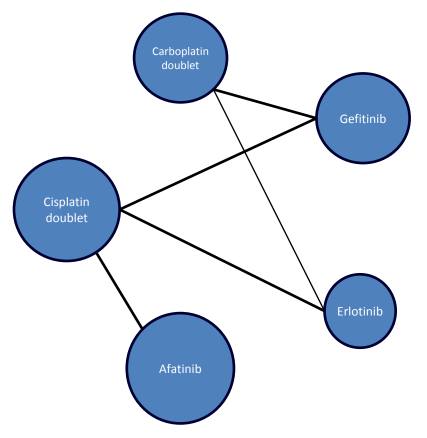


Table D3. Overall survival studies of TKIs (cisplatin-based and carboplatin-based doublets as separate groups)

Study name	Treatment 1	Treatment 2	n1	n2	LogHR	LogSE
LUX LUNG 3	afatinib	cisplatin doublets	230	115	-0.12783	0.146051
LUX LUNG 6	afatinib	cisplatin doublets	242	122	-0.07257	0.134529
IPASS	gefitinib	carboplatin doublets	132	129	0	0.142759
NEJ002	gefitinib	carboplatin doublets	114	114	-0.11991	0.171333
WJTOG3405	gefitinib	cisplatin doublets	86	86	0.224742	0.17812
FIRST-SIGNAL	gefitinib	cisplatin doublets	26	16	0.042101	0.376887
ENSURE	erlotinib	cisplatin doublets	110	107	-0.09	0.19
OPTIMAL	erlotinib	carboplatin doublets	82	72	0.17	0.18
TORCH	erlotinib	cisplatin doublets	19	20	0.46	0.42

Table D4. Network meta-analysis of TKIs: Overall survival (cisplatin-based and carboplatin-based doublets as separate groups) RE Model:

afatinib		_	RE Model: resdev, 7.64 vs. DIC = 0.833	9;
0.91 (0.64 to 1.29)	Cisplatin doublet			
0.83 (0.47 to 1.45)	0.92 (0.58 to 1.42)	Carboplatin doublet		
0.83 (0.49 to 1.40)	0.92 (0.62 to 1.36)	1.00 (0.72 to 1.41)	gefitinib	
0.79 (0.45 to 1.33)	0.87 (0.58 to 1.29)	0.95 (0.60 to 1.47)	0.95 (0.59 to 1.48)	erlotinib

Figure D3. Progression-free survival network diagram of TKIs (all platinum doublets combined)

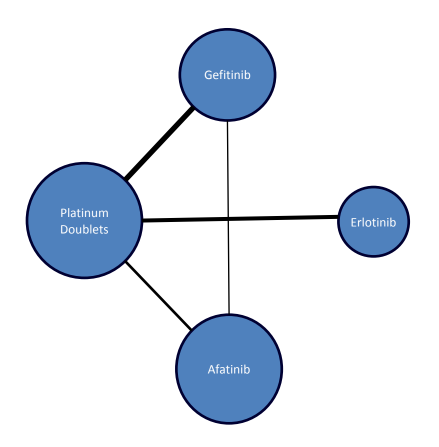


Table D5.	Progression-free surviva	l studies of TKIs (all	platinum doublets combined)

Study name	Treatment 1	Treatment 2	n1	n2	LogHR	LogSE
LUX LUNG 3	afatinib	platinum doublets	230	115	-0.54	0.15
LUX LUNG 6	afatinib	platinum doublets	242	122	-1.27	0.17
LUX LUNG 7	afatinib	gefitinib	160	159	-0.31	0.13
IPASS	gefitinib	platinum doublets	132	129	-0.73	0.15
NEJ002	gefitinib	platinum doublets	114	110	-1.20	0.16
WJTOG3405	gefitinib	platinum doublets	86	86	-0.72	0.19
FIRST-SIGNAL	gefitinib	platinum doublets	26	16	-0.61	0.36
ENSURE	erlotinib	platinum doublets	110	107	-1.08	0.21
OPTIMAL	erlotinib	platinum doublets	82	72	-1.83	0.24
TORCH	erlotinib	platinum doublets	19	20	-0.54	0.38

Table D6. Network meta-analysis of TKIs: Progression-free survival (all platinum doublets combined)

erlotinib		_	RE Model: resdev, 10.53 vs. 10;	
0.79 (0.33 to 1.96)	afatinib		DIC = 5.55	
0.66 (0.30 to 1.50)	0.84 (0.42 to 1.64)	gefitinib		
0.30 (0.16 to 0.58)	0.38 (0.20 to 0.70)	0.45 (0.28 to 0.74)	platinum doublet	

Figure D4. Progression-free survival network diagram of TKIs (cisplatin-based and carboplatin-based doublets as separate groups)

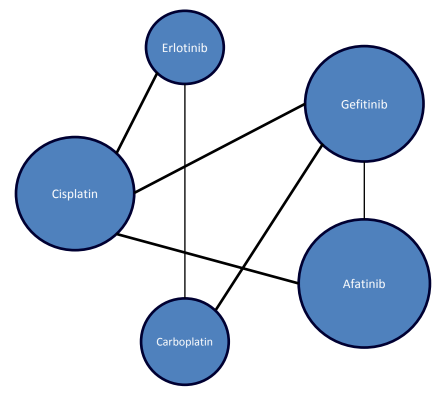


Table D7. Progression-free survival studies of TKIs (cisplatin-based and carboplatin-baseddoublets as separate groups)

Study name	Treatment 1	Treatment 2	n1	n2	LogHR	LogSE
LUX LUNG 3	afatinib	cisplatin doublets	230	115	-0.54	0.15
LUX LUNG 6	afatinib	cisplatin doublets	242	122	-1.27	0.17
LUX LUNG 7	afatinib	gefitinib	160	159	-0.31	0.13
IPASS	gefitinib	carboplatin doublets	132	129	-0.73	0.15
NEJ002	gefitinib	carboplatin doublets	114	110	-1.20	0.16
WJTOG3405	gefitinib	cisplatin doublets	86	86	-0.72	0.19
FIRST-SIGNAL	gefitinib	cisplatin doublets	26	16	-0.61	0.36
ENSURE	erlotinib	cisplatin doublets	110	107	-1.08	0.21
OPTIMAL	erlotinib	Carboplatin doublets	82	72	-1.83	0.24
TORCH	erlotinib	cisplatin doublets	19	20	-0.54	0.38

Table D8. Network meta-analysis of TKIs: Progression-free survival (cisplatin-based and carboplatin-based doublets as separate groups)

erlotinib			RE Model: resdev, 10.19 v DIC = 5.092	s. 10;
0.86 (0.39 to 2.00)	afatinib			
0.62 (0.30 to 1.31)	0.71 (0.37 to 1.34)	gefitinib		
0.36 (0.19 to 0.69)	0.41 (0.24 to 0.72)	0.58 (0.34 to 1.01)	cisplatin	
0.21 (0.10 to 0.44)	0.24 (0.11 to 0.54)	0.34 (0.19 to 0.60)	0.59 (0.28 to 1.17)	carboplatin

Figure D5. Overall survival and progression-free survival network diagram of PD-1 immunotherapies

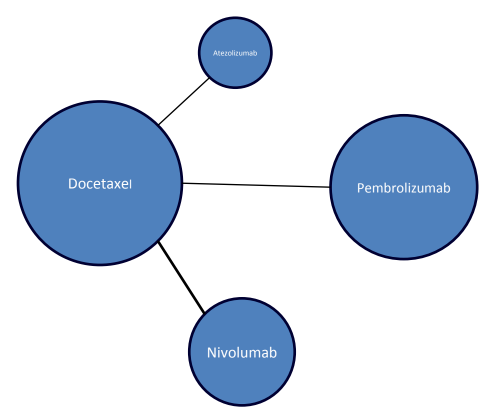


Table D9. Overall survival studies of PD-1 immunotherapies

Study name	Treatment 1	Treatment 2	n1	n2	LogHR	LogSE
CheckMate 017	nivolumab	docetaxel	135	137	-0.52763	0.149301
CheckMate 057	nivolumab	docetaxel	168	172	-0.41552	0.133296
POPLAR	atezolizumab	docetaxel	144	143	-0.31471	0.159395
KEYNOTE-010	pembrolizumab	docetaxel	581*	294	-0.41552	0.095585

*pembrolizumab 2mg/kg and 10mg/kg combined

Nivolumab			FE Model: resdev, 3.3 vs. 4;	
0.95 (0.73 to 1.25)	Pembrolizumab		DIC = -2.564	
0.86 (0.60 to 1.24)	0.90 (0.63 to 1.30)	Atezolizumab		
0.63 (0.52 to 0.76)	0.66 (0.55 to 0.80)	0.73 (0.53 to 1.00)	Docetaxel	

Table D10. Network meta-analysis of PD-1 immunotherapies: Overall survival

Table D11. Progression-free survival	studies of PD-1 immunotherapies
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Study name	Treatment 1	Treatment 2	n1	n2	LogHR	LogSE
CheckMate 017	nivolumab	docetaxel	135	137	-0.47804	0.138852
CheckMate 057	nivolumab	docetaxel	168	172	-0.18633	0.124758
POPLAR	atezolizumab	docetaxel	144	143	-0.06188	0.136612
KEYNOTE-010	pembrolizumab	docetaxel	581*	294	-0.18633	0.082216

*pembrolizumab 2mg/kg and 10mg/kg combined

Nivolumab		res	FE Model: resdev, 5.428 vs. 4; DIC = -1.324				
0.88 (0.69 to 1.12)	Pembrolizumab						
0.77 (0.56 to 1.07)	0.88 (0.65 to 1.21)	Atezolizumab					
0.73 (0.61 to 0.87)	0.83 (0.71 to 0.97)	0.94 (0.72 to 1.23)	Docetaxel				

OS HR	6 months			12 mont	2 months 24 month			hs 60 months				
ref=docetaxel	median	LL	UL	median	LL	UL	median	LL	UL	median	LL	UL
nivolumab	0.65	0.30	1.39	0.52	0.21	1.29	0.42	0.14	1.20	0.32	0.09	1.08
pembrolizumab	0.64	0.15	2.65	0.64	0.12	3.50	0.65	0.09	4.61	0.65	0.06	6.65
atezolizumab	0.70	0.17	2.87	0.57	0.11	3.07	0.47	0.07	3.29	0.37	0.04	3.60

Table D13. Network meta-analysis of parametric OS curves: hazard ratios (Weibull distribution)

Figure D6. Network meta-analysis of parametric OS curves: hazard ratios (Weibull distribution)

Blue: nivolumab Red: pembrolizumab Green: atezolizumab Orange: docetaxel

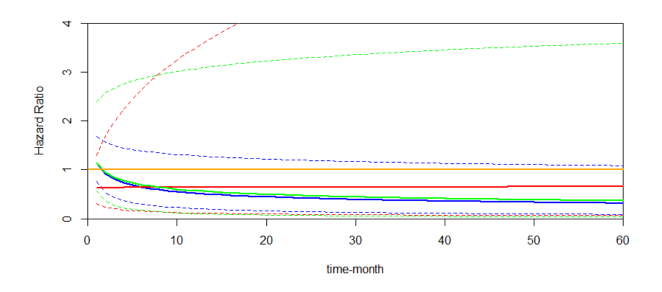


Table D14. Network meta-analysis of parametric OS curves: proportions of OS (Weibull distribution)

Proportion of OS	6 months	12 months	24 months	60 months
	median	median	median	median
ref=docetaxel	61.7%	34.5%	9.5%	0.1%
nivolumab	64.9%	46.4%	25.5%	5.4%
pembrolizumab	73.6%	50.6%	22.1%	1.3%
atezolizumab	64.1%	44.5%	22.9%	3.9%

Figure D7. Network meta-analysis of parametric OS curves: proportions of OS (Weibull distribution)

Blue: nivolumab Red: pembrolizumab Green: atezolizumab Orange: docetaxel

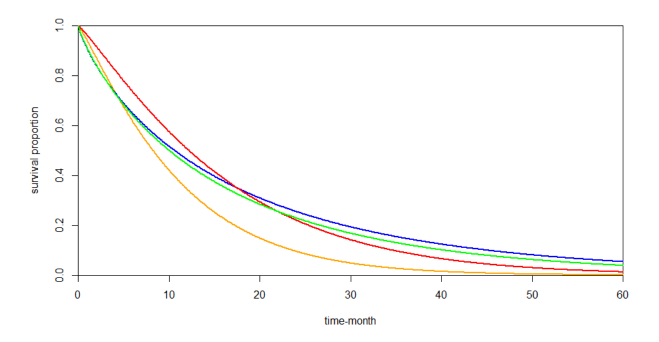


Table D15. Network meta-analysis of parametric PFS curves: hazard ratios (Gompartzdistribution)

PFS HR	1 month	1	3 months		6 months		9 months			12 months					
ref=docetaxel	median	LL	UL	median	LL	UL	median	LL	UL	median	LL	UL	median	LL	UL
nivolumab	1.17	0.85	1.61	0.87	0.57	1.33	0.56	0.31	0.99	0.36	0.17	0.74	0.23	0.09	0.56
pembrolizumab	0.71	0.38	1.30	0.60	0.27	1.32	0.47	0.16	1.34	0.37	0.10	1.36	0.28	0.06	1.39
atezolizumab	1.09	0.65	1.86	0.97	0.49	1.98	0.83	0.31	2.17	0.70	0.20	2.38	0.59	0.13	2.61

Figure D8. Network meta-analysis of parametric PFS curves: hazard ratios (Gompartz distribution)

Blue: nivolumab Red: pembrolizumab Green: atezolizumab Orange: docetaxel

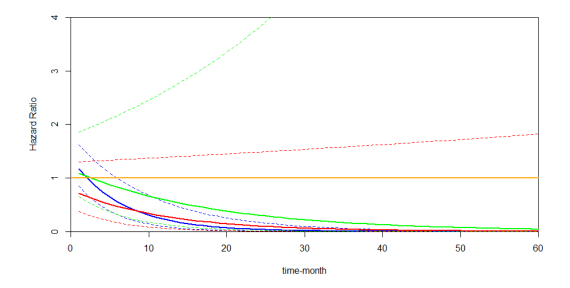
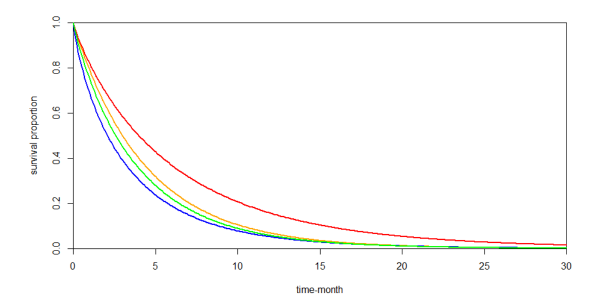


Table D16. Network meta-analysis of parametric PFS curves: proportions of PFS (Gompartz
distribution)

Proportion of PFS	1 months	3 months	6 months	9 months	12 months
	median	median	median	median	median
ref=docetaxel	78.9%	50.0%	25.5%	13.1%	6.7%
nivolumab	68.6%	39.0%	18.8%	9.6%	5.1%
pembrolizumab	81.8%	58.5%	36.8%	23.8%	15.6%
atezolizumab	74.9%	45.1%	22.1%	11.1%	5.7%

Figure D9. Network meta-analysis of parametric PFS curves: proportions of PFS (Gompartz distribution)

Blue: nivolumab Red: pembrolizumab Green: atezolizumab Orange: docetaxel



Appendix E. Subgroup Meta-Analysis Methods and Results

Figure E1. Overall meta-analysis of TKIs: Overall survival

l²=5.036. P=0.391

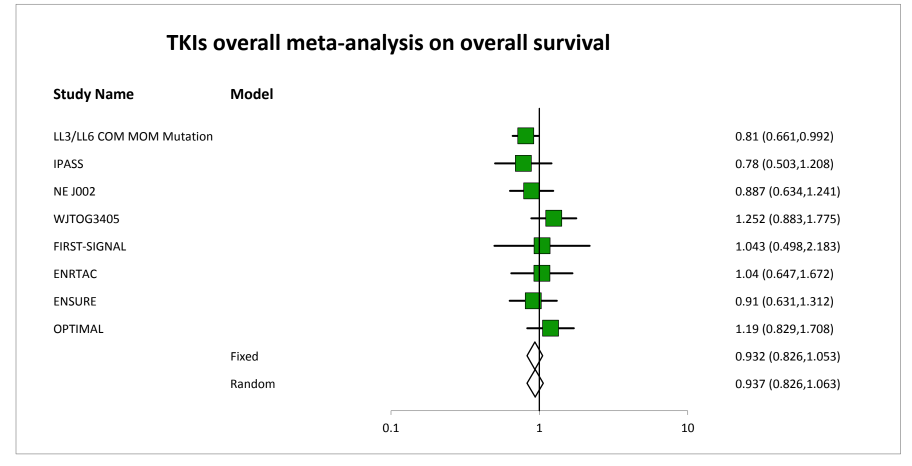


Figure E2. Subgroup meta-analysis of TKIs: Overall survival by mutation type

Mixed Effects Analysis: P=0.115

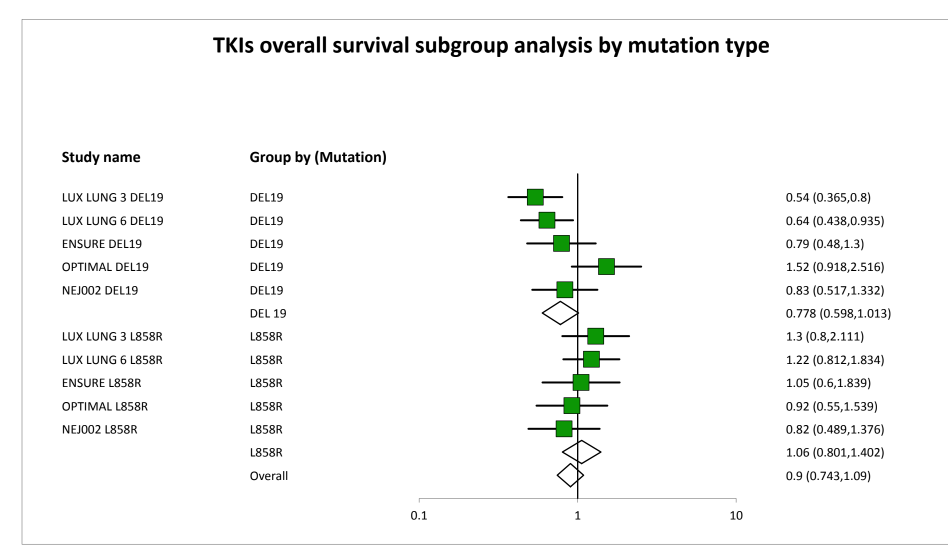


Figure E3. Overall meta-analysis of TKIs: Progression-free survival

l²=70.285. P<0.0001

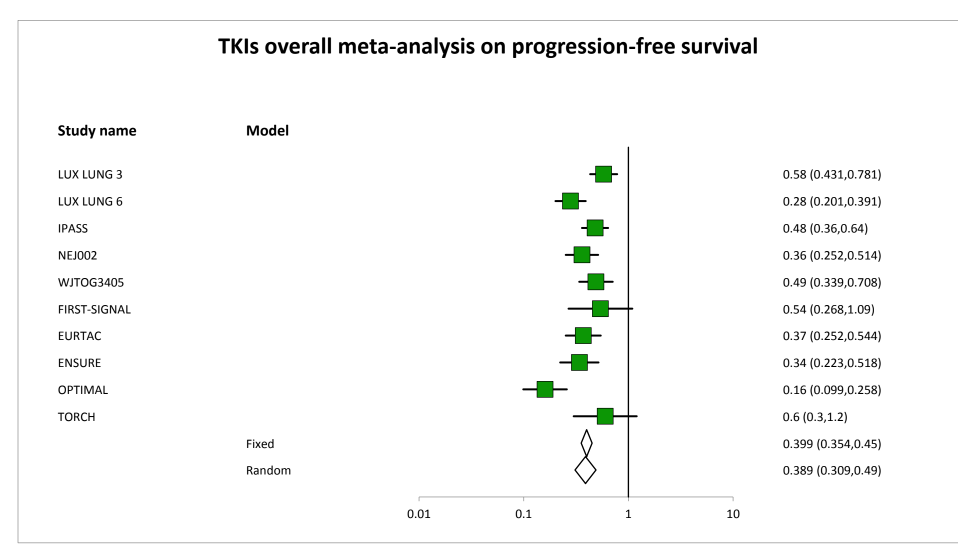


Figure E4. Subgroup meta-analysis of TKIs: Progression-free survival by mutation type

Mixed Effects Analysis: P=0.004

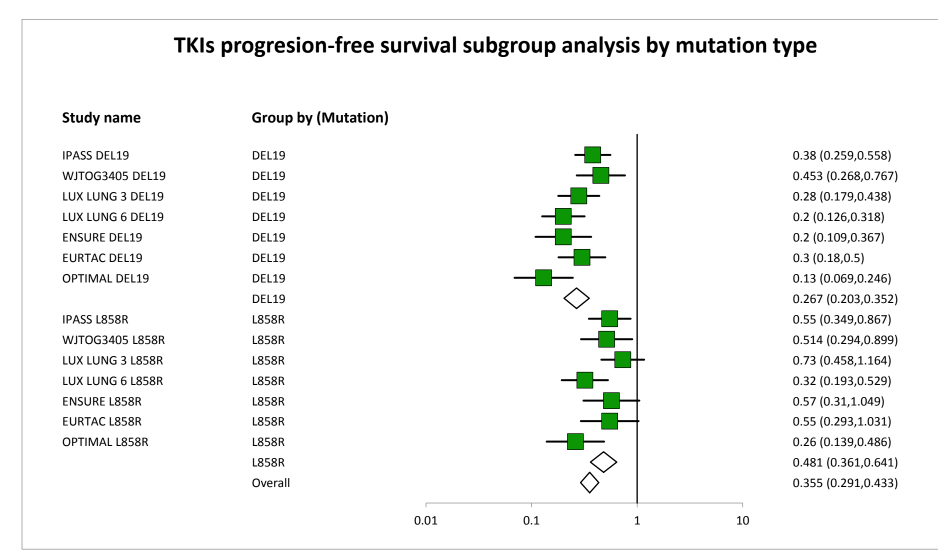


Figure E5. Overall meta-analysis of PD-1 immunotherapies: Overall survival

I²<0.001. P=0.670

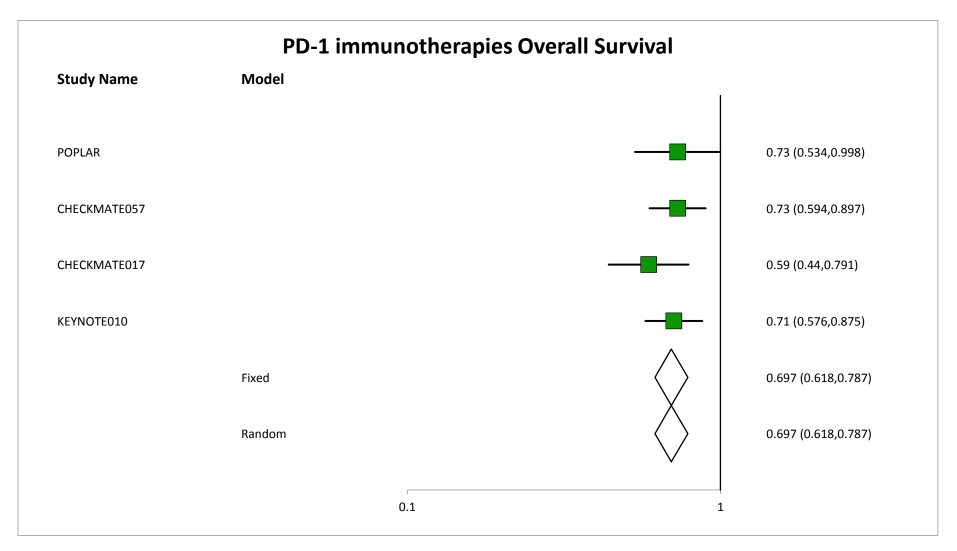


Figure E6. Subgroup meta-analysis of PD-1 immunotherapies: Overall survival by mutation type

Mixed Effects Analysis: P=0.036

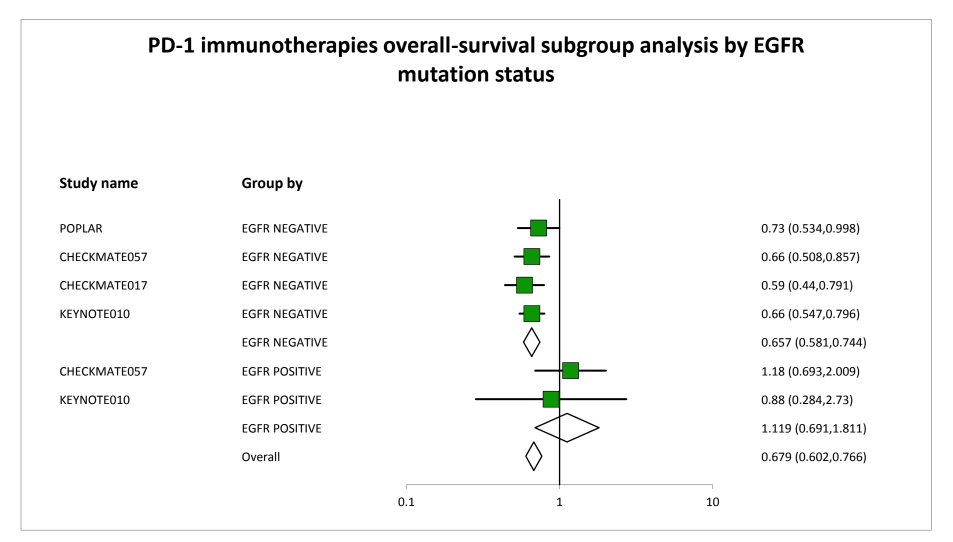


Figure E7. Subgroup meta-analysis of PD-1 immunotherapies: Overall survival by histology

Mixed Effects Analysis: P=0.847

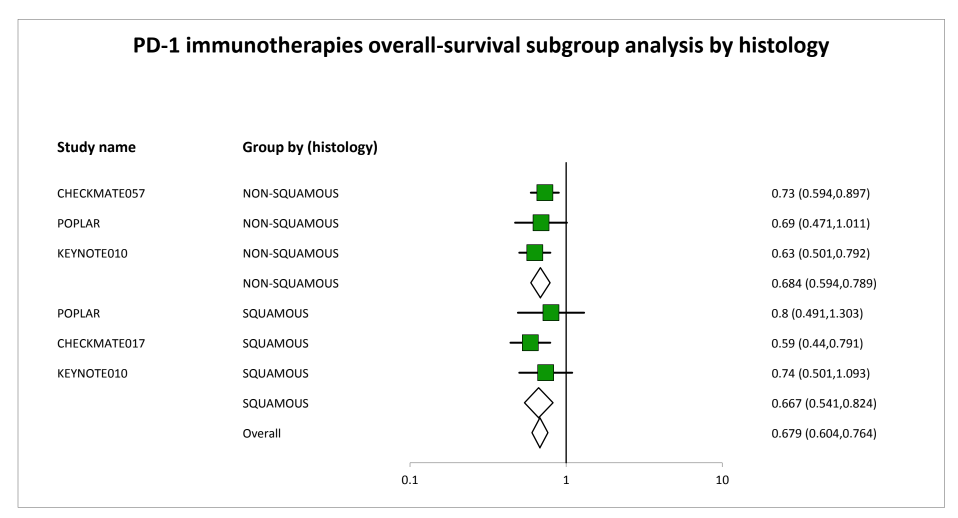


Figure E8. Overall meta-analysis of PD-1 immunotherapies: Progression-free survival

I²=55.485. P=0.081

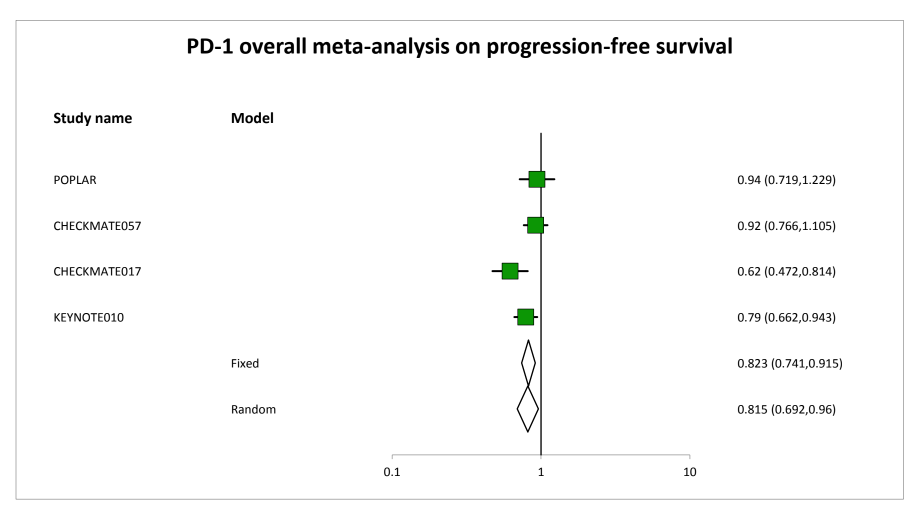


Figure E9. Subgroup meta-analysis of PD-1 immunotherapies: Progression-free survival by mutation type

Mixed Effects Analysis: P=0.002

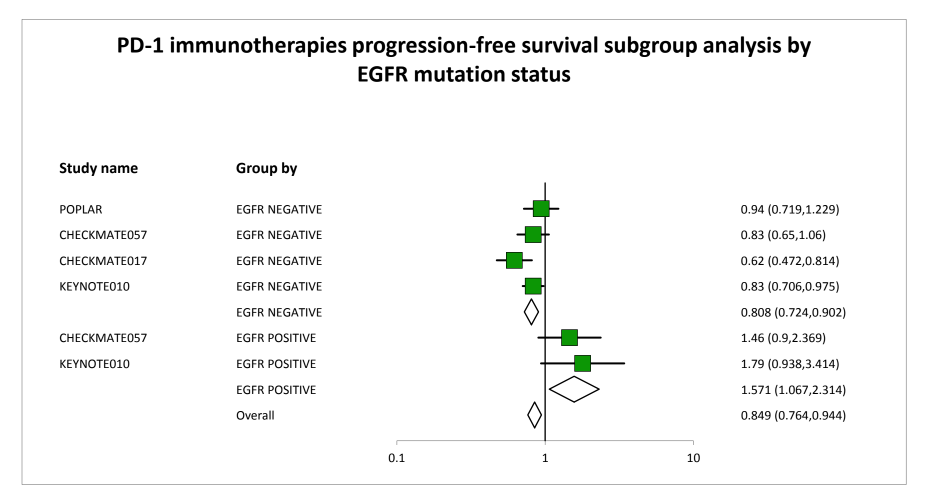
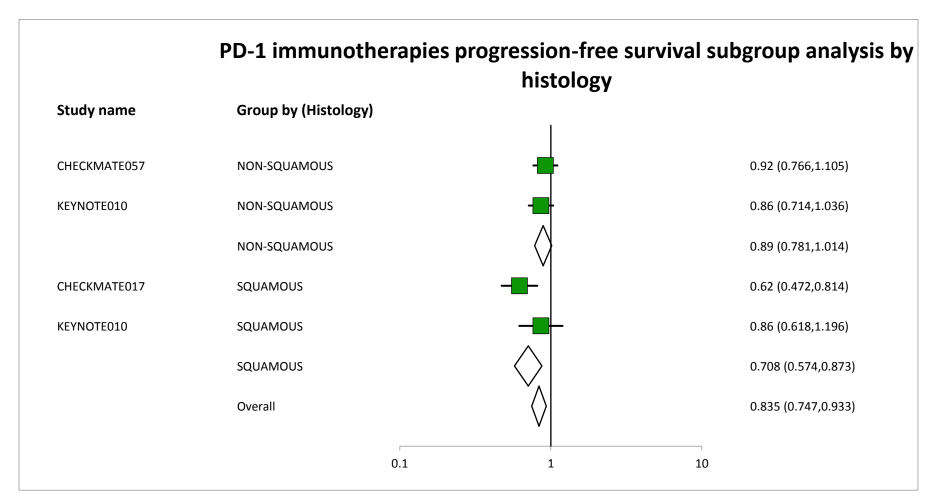


Figure E10. Subgroup meta-analysis of PD-1 immunotherapies: Progression-free survival by histology

Mixed Effects Analysis: P=0.104



Appendix F. Comparative Value Supplemental Information

EGFR+ TKI Therapy	Default	<range></range>		SE	Distribution	Reference
PFS HRs						
AFAT	0.40	0.19	0.82	0.37	LogNormal	NMA
ERLO	0.42	0.18	1.05	0.45	LogNormal	NMA
GEFI	0.53	0.25	1.10	0.38	LogNormal	NMA
OS HRs						
AFAT	0.48	0.38	0.58	0.10	LogNormal	Assumption
ERLO	0.48	0.38	0.58	0.10	LogNormal	Assumption
GEFI	0.48	0.38	0.58	0.10	LogNormal	Assumption

Table F1. Survival hazard ratios in treatment-naïve EGFR patients, from NMA

Table F2. Progression-free survival hazard ratios in second-line immunotherapy patients, fromNMA

Second-Line Immunotherapy	Default	< R:	ange >	Distribu n	ıtio
PFS Hazard Ratios vs. DOCX					
ATEZ: TC1/2/3 or IC1/2/3	0.85	0.63	1.16	6 LogNor	ma
NIVO: All Comers	0.77	0.52	1.13	3 LogNor	ma
PEMB: PD-L1 >1%	0.88	0.74	1.0	5 LogNor	ma
OS Hazard Ratios vs. DOCX					
ATEZ: TC1/2/3 or IC1/2/3	0.59	0.40	0.8	5 LogNor	ma
NIVO: All Comers	0.67	0.55	0.83	3 LogNor	ma
PEMB: PD-L1 >1%	0.71	0.58	0.88	B B LogNor	ma
Second-Line Immunotherapy	Default		< Ran	ge >	Distribution
PFS Hazard Ratios vs. DOCX					
ATEZ: TC1/2/3 or IC1/2/3	0.85	().63	1.16	LogNormal
	0.77	(0.52	1.13	LogNormal
PEMB: PD-L1 >1%	0.88	0).74	1.05	LogNormal
OS Hazard Ratios vs. DOCX					
	0.59	C	0.40	0.85	LogNormal
NIVO: All Comers	0.67	0).55	0.83	LogNormal
	0.71	().58	0.88	LogNormal

Model Survival Curve Fitting

The candidate model curves included the distributional forms Weibull, exponential, log-normal, and log-logistic. We selected the Weibull parametric function in the base case based on face validity, the Akaike Information Criterion (AIC), a graphical assessment of each parametric function, and a knowledge of the expected extrapolation of the progression free survival times (Table 2). Low values for AIC indicate a better mathematical assessment of the fit of the parametric function to the data. It is of note that, while log-logistic and log-normal distributions had generally lower AIC values than the Weibull curves, the tendencies of the former functions to extrapolate long tails was evident on inspection and these distributions were thus ruled out.

Base case PFS and OS curves for CIS-PEM were derived from parametric fits to Kaplan-Meier data from a phase III, non-inferiority, randomized study of cisplatin plus gemcitabine compared with CIS-PEM in chemotherapy-naïve patients with advanced-stage NSCLC.² Base case PFS and OS curves for DOCX were derived from digitized curves from the trials.

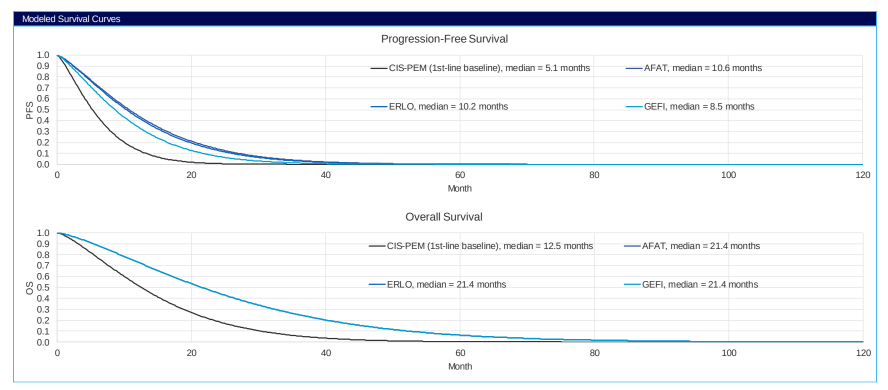
We then used PFS and OS hazard ratios acquired from the network meta-analysis, applied to the universal comparator curves, to derive survival curves for the other interventions (Tables F3 & F4). This approach allowed us to model the relative efficacy of the interventions and survival beyond available follow-up time. We assumed that the treatment effects followed a constant proportional hazard effect in the first-line settings, but that PFS and OS hazard ratios varied with time in the second-line setting, to account for observed trial outcomes showing patients on PD-1 and PD-L1 inhibitors who survive exhibit notable "flattening" of survival curves, representing possibly curative treatment (Table F4).

	AIC	lambda	gamma
PFS Curve Parameters, C	IS-PEM		
Exponential	4042.0	0.1551	
Weibull	3960.9	0.0801	1.3006
Log-Logistic	3854.3	0.0250	2.3021
Log-Normal	3878.1	1.5825	-0.0014
OS Curve Parameters, Cl	S-PEM	·	
Exponential	4915.9	0.0635	
Weibull	4829.7	0.0227	1.3553
Log-Logistic	4780.3	0.0085	1.9686
Log-Normal	4760.8	2.4204	0.8574
PFS Curve Parameters, D	OCX	·	
Exponential	453.1	0.2212	
Weibull	455.0	0.2066	1.0351
Log-Logistic	466.9	0.1678	1.6210
Log-Normal	462.5	1.1010	0.6169
OS Curve Parameters, DC	DCX		
Exponential	654.4	0.0935	
Weibull	645.6	0.0404	1.3082
Log-Logistic	656.4	0.0167	1.9541
Log-Normal	656.1	2.0416	0.8878

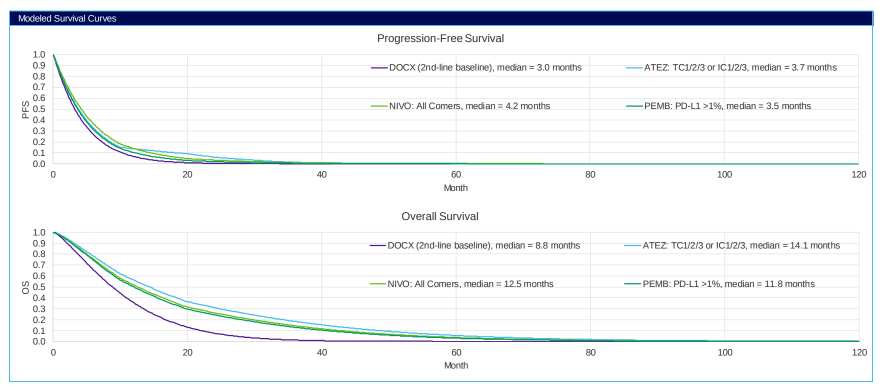
Table F3. Distribution parameters for parametric survival curve fits

Modeled Survival Curves

Figure F1a. 1st-Line Survival: EGFR+







Model Inputs

Table F4. Treatment Regimens and Dosing

	Dosage	Schedule	Route	Duration
Cisplatin	75 mg/m ²	1x/cycle	IV	6 21-day cycles
Pemetrexed	500 mg/m ²	1x/cycle	IV	6 21-day cycles
Docetaxel	75 mg/m ²	every 3 weeks	IV	until progression
Afatinib	40 mg	1x daily	oral	until progression
Erlotinib	150 mg	1x daily	oral	until progression
Gefitinib	250 mg	1x daily	oral	until progression
Atezolizumab	1200 mg	every 3 weeks	IV	until progression
Nivolumab	3 mg/kg	every 2 weeks	IV	until progression
Pembrolizumab	2 mg/kg	every 3 weeks	IV	until progression

Table F5. Other Model Cost Parameters

Other Costs	Default	< Range	>	Distribution
PD-L1 Assay Cost	\$274	\$219	\$329	Normal
End of Life Cost	\$50,000	\$0	\$173,745	Gamma
Prog Supportive Care Cost	\$108.55	\$100.60	\$116.53	Normal
PFS Supportive Care Cost	\$360.48	\$187.57	\$535.50	Normal
TKI Time in Progression (months)	12.0	9.6	14.4	Normal
PD-1 Time in Progression (months)	3.0	2.4	3.6	Normal
Cost of Gemcitabine, per mg	\$0.05	\$0.04	\$0.06	Normal

Table F6. Adverse Event Costs

Cost Per Adverse Event	Def	fault	Range		D	istribution	Short name	Reference
Anemia	\$12,110	\$9,688	\$14,532	Normal	ae_cost_anemia			DRG 808
Diarrhea	\$6,462	\$5,170	\$7,755	Normal	ae_cost_diarrhea			DRG 391
Dyspnea	\$3,951	\$3,161	\$4,741	Normal	ae_cost_dyspnea			DRG 204
Fatigue	\$6,136	\$4,909	\$7,363	Normal	ae_cost_fatigue			DRG 947
Hyponatremia	\$6,133	\$4,907	\$7,360	Normal	ae_cost_hypona			DRG 640
Infection	\$10,314	\$8,251	\$12,377	Normal	ae_cost_infection			DRG 177
Leukopenia	\$12,110	\$9,688	\$14,532	Normal	ae_cost_leukop			DRG 808
Nausea	\$6,462	\$5,170	\$7,755	Normal	ae_cost_nausea			DRG 391
Neuromotor	\$6,926	\$5,541	\$8,311	Normal	ae_cost_neurom			DRG 945
Neutropenia	\$12,110	\$9,688	\$14,532	Normal	ae_cost_neutrop			DRG 808
Paronychia/Nail disorders	\$7,788	\$6 <i>,</i> 230	\$9,345	Normal	ae_cost_paronych			DRG 602
Pneumonitis/Pneumonia	\$7,728	\$6,183	\$9,274	Normal	ae_cost_pneumo			DRG 193
Pulmonary/Respiratory Infx.	\$10,314	\$8,251	\$12,377	Normal	ae_cost_respinfx			DRG 177
Rash	\$7,429	\$5,943	\$8,914	Normal	ae_cost_rash			DRG 606
Skin reactions	\$7,429	\$5,943	\$8,914	Normal	ae_cost_skinrxn			DRG 606
Stomatitis	\$8,101	\$6,481	\$9,721	Normal	ae_cost_stomat			DRG 157

Table F7. Health state utilities

Quality of Life Parameters	Default	< Range >		Distribution	Short Name	Reference
1L PF disease 1L Progressed disease	0.78 0.67	0.77 0.59	0.80 0.75	Beta Beta	util_pf_1L util_prog_1L	LUX-Lung Chouaid et al.
TE FTOGIESSEU UISEASE	0.07	0.59	0.75	Dela	utii_prog_ rL	Chouaid et al.
2L PF disease	0.65	0.61	0.70	Beta	util_pf_2L	Nafees et al.
2L Progressed disease	0.47	0.43	0.52	Beta	util_prog_2L	Nafees et al.
Anemia	0.090	0.059	0.120	Beta	disu anemia	Nafees et al.
Diarrhea	0.090	0.059	0.120	Beta	disu_anernia disu_diarrhea	Nafees et al.
Dyspnea	0.050	0.026	0.074	Beta	disu dyspnea	Doyle et al.
Fatigue	0.073	0.037	0.110	Beta	disu_fatigue	Nafees et al.
Hyponatremia	0.090	0.059	0.120	Beta	disu hyponat	Nafees et al.
Infection	0.047	0.016	0.077	Beta	disu_infex	Nafees et al.
Leukopenia	0.090	0.059	0.120	Beta	disu_leukop	Nafees et al.
Nausea	0.048	0.016	0.080	Beta	disu_nausea	Nafees et al.
Neuromotor	0.069	0.045	0.093	Beta	disu_nueromo	Doyle et al.
Neutropenia	0.090	0.059	0.120	Beta	disu_neutrop	Nafees et al.
Paronychia/Nail disorders	0.032	0.010	0.055	Beta	disu_paronyc	Nafees et al.
Pneumonitis/Pneumonia	0.073	0.037	0.110	Beta	disu_pneumo	Nafees et al.
Pulmonary/Respiratory Infx.	0.046	0.024	0.068	Beta	disu_pulminx	Doyle et al.
Rash	0.032	0.010	0.055	Beta	disu_rash	Nafees et al.
Skin reactions	0.032	0.010	0.055	Beta	disu_skinrx	Nafees et al.
Stomatitis	0.032	0.010	0.055	Beta	disu_stomat	Nafees et al.

PD-1 HRs for Scenarios	Mean	Lower	Uppe r	Mean	Lower	Upper	Mean	Lower	Uppe r	Mean	Lower	Uppe r	Mean	Lower	Uppe r	Mean	Lower	Uppe r
PFS HRs		SE SE		SC	EN: ATE2 Comers	z Ali		: ATEZ 0%			l: NIVO 1%			I: NIVO 0%			: PEMB 50%	
ATEZ	0.85	0.63	1.1 6	0.94	0.72	1.23	0.60	0.31	1.1 6									
NIVO	0.77	0.52	1.1 3							0.69	0.55	0.8 7	0.54	0.40	0.7 2			
PEMB	0.88	0.74	1.0 5													0.59	0.45	0.7 8
OS HRs		SE SE		SC	EN: ATE2 Comers			: ATEZ 0%			l: NIVO 1%			I: NIVO 0%			: PEMB 50%	
ATEZ	0.59	0.40	0.8 5	0.73	0.53	0.99	0.49	0.22	1.0 7									
NIVO	0.67	0.55	0.8 3							0.61	0.48	0.7 9	0.43	0.31	0.5 9			
PEMB	0.71	0.58	0.8 8													0.53	0.40	0.7 0

Table F8. Scenario Analysis 4-8 Hazard Ratios

PSA Results

Table F9. Base case results: TKIs

Results by Regimen								
	CIS-PEM		AFAT		ERLO		GEFI	
TKI Therapy, 1st-Line	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range
Total Costs	\$135,064	(\$60,427 - \$413,243)	\$225,453	(\$121,599 - \$509,683)	\$227,766	(\$117,730 - \$514,988)	\$208,609	(\$109,944 - \$490,759
Drug Costs	\$32,027	(\$25,513 - \$38,338)	\$99,863	(\$52,856 - \$172,455)	\$103,915	(\$48,221 - \$194,061)	\$83,831	(\$43,575 - \$146,901
PFS Supp. Care Costs	\$10,228	(\$9,631 - \$10,789)	\$22,699	(\$12,388 - \$38,623)	\$22,936	(\$10,779 - \$42,937)	\$18,380	(\$9,718 - \$31,958)
Administration Costs	\$1,148	(\$987 - \$1,321)	\$0	\$0	\$0	\$0	\$0	\$0
Progression Costs	\$14,875	(\$11,459 - \$18,685)	\$29,379	(\$1,533 - \$55,565)	\$29,287	(\$31 - \$57,482)	\$36,229	(\$11,390 - \$60,376)
Death Costs	\$71,784	(\$64 - \$349,780)	\$69,773	(\$63 - \$338,987)	\$69,768	(\$63 - \$339,560)	\$69,793	(\$63 - \$340,323)
Adverse Event Costs	\$5,002	(\$3,698 - \$6,559)	\$3,739	(\$2,780 - \$4,846)	\$1,861	(\$1,206 - \$2,686)	\$376	(\$126 - \$754)
Total QALYs	0.88	(0.81 - 0.95)	1.53	(1.24 - 1.88)	1.54	(1.24 - 1.89)	1.50	(1.22 - 1.86)
PFS QALYs	0.42	(0.40 - 0.44)	0.92	(0.51 - 1.53)	0.93	(0.44 - 1.70)	0.75	(0.40 - 1.28)
Progression QALYs	0.46	(0.39 - 0.53)	0.61	(0.03 - 1.13)	0.61	(0.00 - 1.15)	0.76	(0.24 - 1.20)
Total Life Years (OS)	1.22	(1.16 - 1.29)	2.09	(1.67 - 2.56)	2.10	(1.68 - 2.57)	2.08	(1.68 - 2.56)
PFS LYs	0.54	(0.51 - 0.56)	1.17	(0.65 - 1.95)	1.18	(0.57 - 2.16)	0.95	(0.51 - 1.64)
Progression LYs	0.68	(0.62 - 0.76)	0.92	(0.05 - 1.64)	0.91	(0.00 - 1.70)	1.13	(0.37 - 1.77)
Median PFS (months)	5.1	(4.86 - 5.55)	11.6	(6.24 - 19.81)	11.8	(5.55 - 22.11)	9.4	(4.86 - 16.36)
Median OS (months)	12.4	(11.53 - 13.14)	21.7	(17.28 - 26.94)	21.8	(17.28 - 26.94)	21.7	(17.28 - 26.94)

Incremental Results

	CIS-PEM		AFAT		ERLO		GEFI	
TKI Therapy, 1st-Line	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range
ICER			\$138,440	(\$76,088 - \$243,362)	\$140,655	(\$71,337 - \$242,541)	\$117,901	(\$61,935 - \$212,041)
Incremental Costs			\$90,389	(\$47,210 - \$156,737)	\$92,703	(\$41,412 - \$177,254)	\$73,545	(\$34,182 - \$131,714)
Drug Costs			\$67.836	(\$20,808 - \$140,375)	\$71,888	(\$15,277 - \$161,533)	\$51,804	(\$10,605 - \$115,847)
PFS Supp. Care Costs			\$12,472	(\$2,296 - \$28,386)	\$12,708	(\$627 - \$32,683)	\$8,152	(-\$510 - \$21,867)
Administration Costs			-\$1,148	(-\$1,321\$987)	-\$1,148	(-\$1,321\$987)	-\$1,148	(-\$1,321\$987)
Progression Costs			\$14,504	(-\$12,838 - \$39,830)	\$14,412	(-\$14,198 - \$41,966)	\$21,355	(-\$4,216 - \$45,605)
Death Costs			-\$2,011	(-\$10,467\$2)	-\$2,016	(-\$10,362\$1)	-\$1,991	(-\$10,360\$1)
Adverse Event Costs			-\$1,264	(-\$3,021 - \$446)	-\$3,141	(-\$4,876\$1,620)	-\$4,626	(-\$6,233\$3,247)
Incremental QALYs			0.65	(0.38 - 0.98)	0.66	(0.38 - 0.99)	0.62	(0.36 - 0.96)
PFS QALYs			0.50	(0.09 - 1.11)	0.51	(0.03 - 1.27)	0.33	(-0.02 - 0.87)
Progression QALYs			0.16	(-0.41 - 0.64)	0.15	(-0.44 - 0.68)	0.30	(-0.20 - 0.73)
Incremental Life Years (OS)			0.87	(0.46 - 1.32)	0.87	(0.48 - 1.34)	0.86	(0.47 - 1.32)
PFS LYs			0.63	(0.12 - 1.41)	0.64	(0.03 - 1.62)	0.42	(-0.03 - 1.10)
Progression LYs			0.23	(-0.62 - 0.96)	0.23	(-0.66 - 1.02)	0.44	(-0.32 - 1.09)

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Table F10. Base case results: PD-1s

Results by Regimen								
	DOCX		ATEZ: TC1	1/2/3 or IC1/2/3	NIVO: All (Comers	PEMB: PD)-L1 >1%
TKI Therapy, 1st-Line	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range
Total Costs	\$111,451	(\$37,232 - \$394,193)	\$196,693	(\$93,267 - \$464,891)	\$193,707	(\$94,587 - \$476,714)	\$178,589	(\$91,499 - \$458,415
Drug Costs	\$9,610	(\$7,213 - \$12,323)	\$86,216	(\$49,897 - \$126,179)	\$89,220	(\$52,131 - \$145,652)	\$74,686	(\$52,706 - \$104,717
PFS Supp. Care Costs	\$7,278	(\$6,063 - \$8,714)	\$10,769	(\$6,245 - \$15,529)	\$10,897	(\$6,548 - \$17,715)	\$8,920	(\$6,645 - \$11,921)
Administration Costs	\$954	(\$722 - \$1,231)	\$1,251	(\$771 - \$1,987)	\$2,048	(\$1,194 - \$3,375)	\$1,153	(\$799 - \$1,598)
Progression Costs	\$1,856	(\$1,240 - \$2,576)	\$25,161	(\$12,335 - \$45,431)	\$20,071	(\$9,485 - \$31,489)	\$20,771	(\$12,295 - \$31,985)
Death Costs	\$72,525	(\$65 - \$353,826)	\$72,422	(\$62 - \$345,244)	\$71,106	(\$64 - \$348,326)	\$71,273	(\$63 - \$349,911)
Adverse Event Costs	\$19,228	(\$16,405 - \$22,310)	\$874	(\$476 - \$1,424)	\$366	(\$133 - \$708)	\$1,786	(\$1,120 - \$2,615)
Total QALYs	0.48	(0.42 - 0.55)	0.90	(0.62 - 1.26)	0.80	(0.63 - 1.00)	0.75	(0.59 - 0.96)
PFS QALYs	0.25	(0.20 - 0.30)	0.33	(0.21 - 0.52)	0.37	(0.22 - 0.60)	0.30	(0.22 - 0.40)
Progression QALYs	0.24	(0.17 - 0.31)	0.57	(0.27 - 0.94)	0.43	(0.21 - 0.66)	0.45	(0.27 - 0.66)
Total Life Years (OS)	0.88	(0.77 - 1.01)	1.71	(1.14 - 2.48)	1.49	(1.15 - 1.88)	1.42	(1.08 - 1.83)
PFS LYs	0.38	(0.32 - 0.46)	0.51	(0.33 - 0.79)	0.57	(0.34 - 0.90)	0.47	(0.35 - 0.62)
Progression LYs	0.50	(0.36 - 0.64)	1.20	(0.59 - 2.00)	0.92	(0.45 - 1.38)	0.96	(0.58 - 1.37)
Median PFS (months)	3.1	(2.33 - 3.94)	3.8	(2.56 - 5.32)	4.3	(2.56 - 6.47)	3.6	(2.56 - 4.63)
Median OS (months)	8.7	(7.39 - 10.15)	14.5	(9.46 - 21.66)	12.6	(9.46 - 16.13)	12.0	(9.00 - 15.44)

Incremental Results

	DOCX		ATEZ: TC	1/2/3 or IC1/2/3	NIVO: All	Comers	PEMB: PI	D-L1 >1%
TKI Therapy, 1st-Line	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range	Mean	Credible Range
ICER		-	\$200,569	(\$95,341 - \$434,891)	\$256,501	(\$140,016 - \$455,566)	\$248,242	(\$147,003 - \$465,662
Incremental Costs			\$85.242	(\$42,770 - \$126,005)	\$82,256	(\$43,268 - \$137,375)	\$67,138	(\$43,449 - \$96,916)
Drug Costs			\$76,606	(\$41,124 - \$116,026)	\$79,610	(\$43,147 - \$135,361)	\$65,076	(\$43,669 - \$94,498)
PFS Supp. Care Costs			\$3,491	(-\$539 - \$7,744)	\$3,619	(-\$467 - \$10,070)	\$1,642	(-\$72 - \$3,993)
Administration Costs			\$297	(-\$191 - \$965)	\$1,094	(\$258 - \$2,413)	\$199	(-\$157 - \$590)
Progression Costs			\$23,305	(\$10,553 - \$43,618)	\$18,215	(\$7,853 - \$29,652)	\$18,915	(\$10,711 - \$29,862)
Death Costs			-\$103	(-\$11,462\$2)	-\$1,419	(-\$7,464\$1)	-\$1,252	(-\$6,643\$1)
Adverse Event Costs			-\$18,354	(-\$21,485\$15,441)	-\$18,863	(-\$21,965\$16,047)	-\$17,442	(-\$20,575\$14,518
Incremental QALYs			0.41	(0.16 - 0.76)	0.32	(0.18 - 0.49)	0.27	(0.14 - 0.44)
PFS QALYs			0.08	(-0.02 - 0.25)	0.12	(-0.01 - 0.33)	0.06	(0.00 - 0.14)
Progression QALYs			0.33	(0.06 - 0.69)	0.20	(0.00 - 0.39)	0.21	(0.07 - 0.39)
Incremental Life Years (OS)			0.83	(0.30 - 1.56)	0.61	(0.33 - 0.95)	0.54	(0.27 - 0.88)
PFS LYs			0.13	(-0.03 - 0.38)	0.18	(-0.02 - 0.51)	0.08	(0.00 - 0.20)
Progression LYs			0.70	(0.12 - 1.48)	0.42	(0.00 - 0.83)	0.46	(0.16 - 0.82)

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Table F11. Scenario 1a (OS benefit turned off)

	CIS-PEM	AFAT	ERLO	GEFI
TKI Therapy, 1st-Line	Deterministic	Deterministic	Deterministic	Deterministic
Total Costs	\$112,361	\$171,419	\$165,984	\$148,632
Drug Costs	\$32,042	\$91,560	\$91,463	\$76,609
PFS Supp. Care Costs	\$10,217	\$20,757	\$19,988	\$16,696
Administration Costs	\$1,145	\$0	\$0	\$0
Progression Costs	\$15,763	\$7,293	\$4,420	\$6,548
Death Costs	\$48,192	\$48,066	\$48,257	\$48,405
Adverse Event Costs	\$5,002	\$3,744	\$1,855	\$375
Total QALYs	0.88	1.00	0.91	0.82
PFS QALYs	0.42	0.84	0.81	0.68
Progression QALYs	0.46	0.15	0.09	0.14
Total Life Years (OS)	1.22	1.31	1.18	1.08
PFS LYs	0.54	1.08	1.04	0.87
Progression LYs	0.68	0.23	0.14	0.21
Median PFS (months)	5.1	10.6	10.2	8.5
Median OS (months)	12.5	13.4	12.0	10.8

Table F12. Scenario 1b (OS benefit turned off)

	CIS-PEM	AFAT	ERLO	GEFI
TKI Therapy, 1st-Line	Deterministic	Deterministic	Deterministic	Deterministic
ICER		\$495,393	\$1,919,236	-\$627,258
Incremental Costs		\$59,059	\$53,623	\$36,271
Drug Costs		\$59,519	\$59,421	\$44,567
PFS Supp. Care Costs		\$10,540	\$9,771	\$6,479
Administration Costs		-\$1,145	-\$1,145	-\$1,145
Progression Costs		-\$8,470	-\$11,343	-\$9,215
Death Costs		-\$126	\$65	\$213
Adverse Event Costs		-\$1,259	-\$3,147	-\$4,628
Incremental QALYs		0.12	0.03	-0.06
PFS QALYs		0.42	0.39	0.26
Progression QALYs		-0.30	-0.36	-0.32
Incremental Life Years (OS)		0.09	-0.04	-0.14
PFS LYs		0.54	0.50	0.33
Progression LYs		-0.45	-0.54	-0.48

Table F13. Scenario 2: ATEZ All Comers

lesults by Regimen			Incremental Results		
	DOCX	SCEN: ATEZ All Comers		DOCX	SCEN: ATEZ All Comers
Immunotherapy, 2nd-Line	Deterministic	Deterministic	Immunotherapy, 2nd-Line	Deterministic	Deterministic
Total Costs	\$87,831	\$154,840	ICER		\$270,338
Drug Costs	\$9,563	\$75,231			
PFS Supp. Care Costs	\$7,224	\$9,196	Incremental Costs		\$67,008
Administration Costs	\$947	\$1,184	Drug Costs		\$65,668
Progression Costs	\$2,134	\$20,392	PFS Supp. Care Costs		\$1,972
Death Costs	\$48,693	\$47,967	Administration Costs		\$237
Adverse Event Costs	\$19,270	\$870	Progression Costs		\$18,258
			Death Costs		-\$726
Total QALYs	0.48	0.73	Adverse Event Costs		-\$18,400
PFS QALYs	0.24	0.31			
Progression QALYs	0.24	0.42	Incremental QALYs		0.25
			PFS QALYs		0.07
Total Life Years (OS)	0.88	1.37	Progression QALYs		0.18
PFS LYs	0.38	0.48			
Progression LYs	0.50	0.89	Incremental Life Years (OS)		0.48
Median PFS (months)	3.0	3.3	PFS LYs		0.10
Median OS (months)	8.8	11.5	Progression LYs		0.38

Table F14. Scenario 3: ATEZ >50%

lesults by Regimen			Incremental Results		
	DOCX	SCEN: ATEZ >50%		DOCX	SCEN: ATEZ >50%
Immunotherapy, 2nd-Line	Deterministic	Deterministic	Immunotherapy, 2nd-Line	Deterministic	Deterministic
Total Costs	\$87,831	\$218,215	ICER		\$221,845
Drug Costs	\$9,563	\$126,326			
PFS Supp. Care Costs	\$7,224	\$15,903	Incremental Costs		\$130,383
Administration Costs	\$947	\$1,991	Drug Costs		\$116,763
Progression Costs	\$2,134	\$26,183	PFS Supp. Care Costs		\$8,678
Death Costs	\$48,693	\$46,943	Administration Costs		\$1,044
Adverse Event Costs	\$19,270	\$870	Progression Costs		\$24,049
			Death Costs		-\$1,751
Total QALYs	0.48	1.07	Adverse Event Costs		-\$18,400
PFS QALYs	0.24	0.54			
Progression QALYs	0.24	0.53	Incremental QALYs		0.59
			PFS QALYs		0.30
Total Life Years (OS)	0.88	1.96	Progression QALYs		0.30
PFS LYs	0.38	0.82			
Progression LYs	0.50	1.14	Incremental Life Years (OS)		1.08
Median PFS (months)	3.0	5.3	PFS LYs		0.44
Median OS (months)	8.8	16.8	Progression LYs		0.64

Table F15. Scenario 4: NIVO >1%

lesults by Regimen			Incremental Results		
	DOCX	SCEN: NIVO >1%		DOCX	SCEN: NIVO >1%
Immunotherapy, 2nd-Line	Deterministic	Deterministic	Immunotherapy, 2nd-Line	Deterministic	Deterministic
Total Costs	\$87,831	\$178,879	ICER		\$239,258
Drug Costs	\$9,563	\$94,152			
PFS Supp. Care Costs	\$7,224	\$11,502	Incremental Costs		\$91,047
Administration Costs	\$947	\$2,162	Drug Costs		\$84,589
Progression Costs	\$2,134	\$23,107	PFS Supp. Care Costs		\$4,278
Death Costs	\$48,693	\$47,588	Administration Costs		\$1,215
Adverse Event Costs	\$19,270	\$367	Progression Costs		\$20,973
			Death Costs		-\$1,105
Total QALYs	0.48	0.86	Adverse Event Costs		-\$18,902
PFS QALYs	0.24	0.39			
Progression QALYs	0.24	0.47	Incremental QALYs		0.38
			PFS QALYs		0.14
Total Life Years (OS)	0.88	1.60	Progression QALYs		0.24
PFS LYs	0.38	0.60			
Progression LYs	0.50	1.00	Incremental Life Years (OS)		0.72
Median PFS (months)	3.0	4.6	PFS LYs		0.22
Median OS (months)	8.8	13.6	Progression LYs		0.50

Table F16. Scenario 5: NIVO >10%

lesults by Regimen			Incremental Results		
	DOCX	SCEN: NIVO >10%		DOCX	SCEN: NIVO >10%
Immunotherapy, 2nd-Line	Deterministic	Deterministic	Immunotherapy, 2nd-Line	Deterministic	Deterministic
Total Costs	\$87,831	\$222,773	ICER		\$193,076
Drug Costs	\$9,563	\$125,317			
PFS Supp. Care Costs	\$7,224	\$15,501	Incremental Costs		\$134,941
Administration Costs	\$947	\$2,878	Drug Costs		\$115,754
Progression Costs	\$2,134	\$32,288	PFS Supp. Care Costs		\$8,277
Death Costs	\$48,693	\$46,422	Administration Costs		\$1,931
Adverse Event Costs	\$19,270	\$367	Progression Costs		\$30,154
			Death Costs		-\$2,272
Total QALYs	0.48	1.18	Adverse Event Costs		-\$18,902
PFS QALYs	0.24	0.52			
Progression QALYs	0.24	0.66	Incremental QALYs		0.70
			PFS QALYs		0.28
Total Life Years (OS)	0.88	2.20	Progression QALYs		0.42
PFS LYs	0.38	0.80			
Progression LYs	0.50	1.40	Incremental Life Years (OS)		1.32
Median PFS (months)	3.0	5.8	PFS LYs		0.42
Median OS (months)	8.8	18.9	Progression LYs		0.90

Table F17. Scenario 6: PEMB >50%

lesults by Regimen			Incremental Results		
	DOCX	SCEN: PEMB >50%		DOCX	SCEN: PEMB >50%
Immunotherapy, 2nd-Line	Deterministic	Deterministic	Immunotherapy, 2nd-Line	Deterministic	Deterministic
Total Costs	\$87,831	\$202,675	ICER		\$226,523
Drug Costs	\$9,563	\$112,562			
PFS Supp. Care Costs	\$7,224	\$13,844	Incremental Costs		\$114,844
Administration Costs	\$947	\$1,745	Drug Costs		\$102,999
Progression Costs	\$2,134	\$25,545	PFS Supp. Care Costs		\$6,620
Death Costs	\$48,693	\$47,195	Administration Costs		\$798
Adverse Event Costs	\$19,270	\$1,785	Progression Costs		\$23,411
			Death Costs		-\$1,498
Total QALYs	0.48	0.99	Adverse Event Costs		-\$17,485
PFS QALYs	0.24	0.47			
Progression QALYs	0.24	0.52	Incremental QALYs		0.51
			PFS QALYs		0.22
Total Life Years (OS)	0.88	1.83	Progression QALYs		0.29
PFS LYs	0.38	0.72			
Progression LYs	0.50	1.11	Incremental Life Years (OS)		0.94
Median PFS (months)	3.0	5.3	PFS LYs		0.34
Median OS (months)	8.8	15.7	Progression LYs		0.61

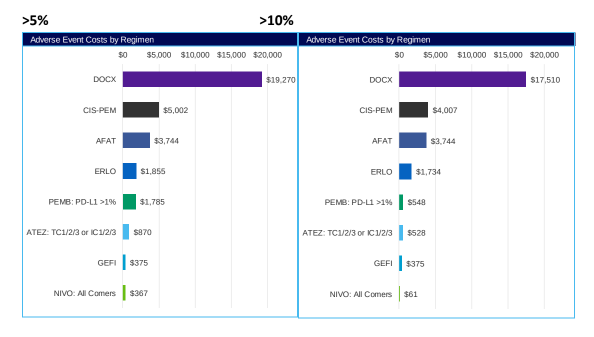


Table F18. Scenario 7: Adverse Event Thresholds

<u>Appendix G. Previous Technology Assessments</u> <u>and Systematic Reviews</u>

We identified four completed technology assessments: three from the National Institute for Health and Care Excellence (NICE) in the UK and one from the Pan-Canadian Oncology Drug Review (pCODR). These reviews of afatinib, erlotinib, and gefitinib are summarized below. Of note, NICE expects to publish a final appraisal document of pembrolizumab in January 2017 and is currently suspending the appeal stage of nivolumab to allow the manufacturer to make a further submission that includes a patient access scheme. We also identified six systematic reviews of the TKIs and a single systematic review of the PD-1 immunotherapies for NSCLC.

Technology Assessments

NICE Gefitinib (2010), Erlotinib (2012), Afatinib (2014)

- Gefitinib: https://www.nice.org.uk/guidance/ta192
- Erlotinib: <u>https://www.nice.org.uk/guidance/ta258</u>
- Afatinib: <u>https://www.nice.org.uk/guidance/ta310</u>

NICE recommends gefitinib, erlotinib, and afatinib for first-line treatment of patients with locally advanced or metastatic NSCLC if they test positive for the epidermal growth factor receptor tyrosine kinase mutation and the manufacturer provides the drugs at a discounted price agreed under the patient access scheme.

Pan-Canadian Oncology Drug Review (pCODR) Expert Review Committee Final Recommendation: Afatinib (Giotrif) for first line treatment of EGFR Mutation Positive Advanced NSCLC (May 2, 2014)

(https://www.cadth.ca/sites/default/files/pcodr/pcodr-giotrif-nsclc-fn-rec.pdf)

The pCODR expert review committee issued a final recommendation (based on the MM-003 trial) that afatinib should be funded if cisplatin-pemetrexed is currently the main treatment option for first-line treatment of EGFR mutation positive advanced or metastatic adenocarcinoma with an ECOG performance status of 0-1. The review committee also recommended funding afatinib as an alternative to gefitinib, provided that cost-effectiveness is improved to an "acceptable" level. Despite the lack of head-to-head evidence, the committee believed afatinib to and gefitinib to provide similar clinical benefit and expressed a commitment to providing access to more treatment options. They noted that afatinib was cost-effective compared to cisplatin-pemetrexed but may not be considered cost-effective relative to gefitinib.

Previous systematic reviews

Several recent systematic reviews and meta-analyses have compared the three EGFR TKIs of interest to one another as well as to platinum-based chemotherapy doublets in chemo-naïve patients with EGFR mutations. These studies have consistently reported superior efficacy with the TKIs relative to chemotherapy, albeit no study has shown improvements in overall survival, which is likely due to high rates of crossover between therapies. Comparisons between TKIs have consistently shown no statistical differences in PFS, ORR, or OS, although toxicity profiles have varied across agents.

Haspinger ER, Agustoni F, Torri V, et al. Is there evidence for different effects among EGFR-TKIs? Systematic review and meta-analysis of EGFR tyrosine kinase inhibitors (TKIs) versus chemotherapy as first-line treatment for patients harboring EGFR mutations. Crit Rev Oncol Hematol. 2015;94:213-227.

Haspinger et al. performed a systematic review and meta-analysis of all phase II/III RCTs published up to June 2014 that examined afatinib, erlotinib, or gefitinib in previously-untreated patients with NSCLC and an EGFR-mutation. The literature search identified 9 RCTs, which involved 1,774 EGFRmutated patients. Direct comparisons of each of the TKIs versus a platinum-based chemotherapy doublet showed statistically significant differences in favor of the TKIs for both PFS and ORR but no difference in overall survival. Indirect comparisons between TKIs did not show statistical differences for PFS, ORR, or OS, although safety profiles varied; afatinib had more events of diarrhea and rash compared to both erlotinib and gefitinib, while gefitinib had a higher rate of hypertransaminasemia. There were no differences in treatment discontinuation and treatmentrelated deaths across agents.

Des Guetz G, Landre T, Uzzan B, et al. Is there a Survival Benefit of First-Line Epidermal Growth Factor Receptor Tyrosine-Kinase Inhibitor Monotherapy Versus Chemotherapy in Patients with Advanced Non-Small-Cell Lung Cancer?: A Meta-Analysis. Targ Oncol. 2016;11:41-47.

A meta-analysis from Des Guetz and colleagues examined eight phase III RCTs of first-line treatment of advanced NSCLC with afatinib, erlotinib, or gefitinib versus a platinum-based chemotherapy doublet. The analysis included 2,962 patients with EGFR mutations. Compared to chemotherapy, TKIs significantly improved PFS (HR 0.37, 95% CI 0.29-0.49) but not overall survival (HR 0.98, 95% CI 0.87-1.10). The authors suggested that a high level of crossover between both groups might explain why the TKIs showed an improved PFS without parallel improvement in OS. Comparisons between TKIs showed no significant differences between gefitinib and erlotinib or afatinib and erlotinib. Grade 3-4 adverse effects differed between TKIs and chemotherapy, with rashes and diarrhea occurring more frequently with TKIs (RR 4.60, 95% CI 2.37-8.95; and RR 3.88, 95% CI 2.00-7.56) and nausea/vomiting (RR 0.22, 95% CI 0.06-0.54), neutropenia (RR 0.06, 95% CI 0.04-0.08), thrombocytopenia (RR 0.11, 95% CI 0.04-0.32), and anemia (RR 0.10, 95% CI 0.04-0.27) occurring more frequently with chemotherapy.

Haaland B, Tan PS, de Castro G, and Lopes G. Meta-Analysis of First-Line Therapies in Advanced Non-Small-Cell Lung Cancer Harboring EGFR-Activating Mutations. J Thorac Oncol. 2014;9:805-811.

Another meta-analysis of gefitinib, erlotinib, afatinib, and chemotherapy in previously untreated patients with advanced non-small-cell lung cancer harboring EGFR-activating mutations reported similar results. Compared to chemotherapy, the analysis showed each of the TKIs to improve PFS and ORR but not OS. Outcomes did not statistically differ between individual TKIs. Similar to the findings of other analyses, diarrhea, rash, and pruritus occurred more frequently with the TKIs, whereas anorexia, anemia, fatigue, nausea, vomiting, alopecia, and neutropenia were more common with chemotherapy.

Zhang Y, Sheng J, Yang Y, et al. Optimized selection of three major EGFR-TKIs in advanced EGFRpositive non-small cell lung cancer: a network meta-analysis. Oncotarget. 2016;7(15):20093-20108.

In a series of analyses of TKI therapy in chemo-naïve and previously treated EGFR+ patients, Zhang and colleagues found similar efficacy among the TKIs for ORR, PFS, and disease control rate relative to chemotherapy. Patients with EGFR exon 19 deletion showed superior numerical data with respect to ORR, 1-year PFS, 1-year OS, and 2-year OS compared with 21 L858R patients. Among the TKIs, afatinib had the highest risk of diarrhea, while gefitinib presented the greatest risk of elevated liver transaminase.

Popat S, Mok T, Yang JC-H, et al. Afatinib in the treatment of EGFR mutation-positive NSCLC-A network meta-analysis. Lung Cancer. 2014;85:230-238.

Popat and colleagues assessed the relative efficacy of afatinib, erlotinib, and gefitinib by conducting a systematic literature review and network meta-analysis; eight of the 21 studies reviewed by the authors reported results from patients with an EGFR+ mutation and were included in the network. Similar to the findings of other analyses, Popat et al. did not find statistical differences in overall survival between treatments. However, this study did suggest that afatinib might have a slight PFS benefit compared to gefitinib (HR 0.60, 95% CI 0.34-0.99) in patients with common EGFR mutations.

Greenhalgh J, Dwan K, Boland A, et al. First-line treatment of advanced epidermal growth factor receptor (EGFR) mutation positive non-squamous non-small cell lung cancer. Cochrane Database of Systematic Reviews. 2016;5.

A review from the Cochrane Collaboration of first-line treatment with afatinib, erlotinib, and gefitinib in non-squamous, EGFR+ NSCLC did not find a statistically-significant overall survival

benefit with single-agent use of the TKIs (vs. chemotherapy), but did find a significant PFS benefit with each of the agents of interest. The most commonly reported grade 3-4 AEs were rash and diarrhea with the TKIs, while myelosuppression, fatigue, and anorexia were common with chemotherapy. Quality of life and symptoms were improved in one or more indices measured in two trials each for the three TKIs of focus.

Melosky B, Chu Q, Juergens R, et al. Pointed Progress in Second-Line Advanced Non-Small-Cell Lung Cancer: The Rapidly Evolving Field of Checkpoint Inhibition. J Clin Oncol. 2016;34:1676-1688.

We identified a single systematic review of the PD-1 immunotherapies. The review included 20 studies of PD-1 immunotherapies in the second-line setting, three of which were RCTS. Melosky and colleagues reviewed the phase II POPLAR trial of atezolizumab as well as the two phase III CheckMate trials of nivolumab in squamous and non-squamous NSCLC, respectively. Relative to docetaxel, atezolizumab showed a trend toward better overall survival, particularly in patients with increased PD-L1 expression. Nivolumab improved overall survival in patients with both squamous and nonsquamous carcinoma, although PD-L1 expression was only associated with overall survival in those with nonsquamous disease. The authors concluded that PD-1 immunotherapy is safe compared to standard chemotherapy and is best suited for patients with higher levels of PD-L1 expression and no EGFR mutations.

Appendix H. Ongoing Studies

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
Afatinib					
A Randomised, Open- label, Phase III Study of BIBW 2992 Versus Chemotherapy as First-line Treatment for Patients with Stage IIIB or IV Adenocarcinoma of the Lung Harbouring an EGFR Activating Mutation NCT00949650	RCT	October 2016	 Afatinib Cisplatin + Pemetrexed 	 N=345 Age 18 years and older Inclusion Criteria: Stage IIIB/IV NSCLC EGFR mutation positive ECOG score of 0-1 Life expectancy of at least 3 months Exclusion Criteria: Prior chemotherapy for relapsed or metastatic NSCLC or EGFR TKIs Radiotherapy within 4 weeks Active brain metastases Any significant illness or organ dysfunction Hep B, Hep C or HIV carrier 	 <u>Primary</u>: PFS [Time Frame: every 12 weeks until death] <u>Secondary</u>: ORR OS DC QoL ECOG PS

Title/Trial Name	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
Gefitinib vs. Erlotinib					
A Randomized, Open- label Phase II Trial of Erlotinib 100mg Daily Versus Gefitinib 250mg Daily in Patients with Advanced Non-Small Cell Lung Cancer Who Harbor EGFR Mutations. NCT01955421	RCT	June 2016	 Erlotinib 100mg Gefitinib 250mg 	 N=224 Age 18 years and older <u>Inclusion Criteria</u>: Stage IIIB or IV NSCLC EGFR mutation, including exon 19 or exon 21 L858R Measurable disease according to RECIST 1.1 ECOG PS 0-2 Adequate organ function <u>Exclusion Criteria</u>: Prior treatment with EGFR TKIs Radiotherapy within 4 weeks Active brain metastases Any other current malignancy or malignancy diagnosed within in past 3 years 	 <u>Primary</u>: Disease control rate [Time Frame: 2 years] <u>Secondary</u>: PFS AEs

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
Nivolumab					
An Open-Label, Randomized Phase 3 Trial of Nivolumab, or Nivolumab Plus Ipilimumab, or Nivolumab Plus Platinum Doublet Chemotherapy Versus Platinum Doublet Chemotherapy in Subjects with Chemotherapy-Naïve Stage IV or Recurrent Non-Small Cell Lung Cancer (NSCLC) NCT02477826	RCT	December 2020	 Nivolumab Nivolumab + Ipilimumab Nivolumab + Platinum doublet chemotherapy Platinum doublet chemotherapy 	 N=1980 Age 18 years and older Inclusion Criteria: Stage IV or recurrent NSCLC No prior systemic anti-cancer therapy Have PD-L1 immunohistochemical testing ECOG PS 0-1 Measurable disease according to RECIST 1.1 Exclusion Criteria: CNS metastases Suspected autoimmune diseases Hep B, Hep C or HIV positive 	 <u>Primary</u>: OS, PFS [Time Frame: 48 months] <u>Secondary</u>: ORR Symptoms

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
An Open-label Randomized Multinational Phase 3 Trial of Nivolumab Versus Docetaxel in Previously Treated Subjects with Advanced or Metastatic Non-Small Cell Lung Cancer (CheckMate 078: Checkpoint Pathway and nivolumab Clinica I Trial Evaluation 078) NCT02613507	RCT	January 2019	 Nivolumab Docetaxel 	 N=500 Age 18 years and older Inclusion Criteria: Stage IIIB/IV or recurrent NSCLC Disease progression during or after one prior platinum containing doublet chemotherapy ECOG PS 0-1 Measurable disease according to RECIST 1.1 Exclusion Criteria: CNS metastases Suspected autoimmune diseases Prior docetaxel or checkpoint inhibitors 	 <u>Primary:</u> OS [Time Frame: 37 months] <u>Secondary:</u> ORR PFS Symptoms

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
An Open-Label, Randomized, Phase 3 Trial of Nivolumab Versus Investigator's Choice Chemotherapy as First-Line Therapy for Stage IV or Recurrent PD-L1+ Non-Small Cell Lung Cancer (CheckMate 026) NCT02041533	RCT	January 2018	 Nivolumab Physician's choice chemotherapy 	 N=535 Age 18 years and older Inclusion Criteria: Stage IV or recurrent NSCLC No prior systemic anticancer therapy ECOG PS 0-1 Measurable disease according to RECIST 1.1 Exclusion Criteria: EGFR mutation or ALK translocation CNS metastases Suspected autoimmune diseases 	 Primary: PFS [Time Frame: 33 months] Secondary: ORR PFS Symptoms
An Open-Label Randomized Phase III Trial of BMS- 936558 (Nivolumab) Versus Docetaxel in Previously Treated Advanced or Metastatic Squamous Cell Non- Small Cell Lung Cancer (NSCLC) NCT01642004	RCT	January 2017	 Nivolumab Docetaxel 	 N=352 Age 18 years and older Inclusion Criteria: Stage III/IV or recurrent NSCLC progression during/after one prior platinum doublet-based chemotherapy ECOG PS 0-1 Measurable disease according to RECIST 1.1 Exclusion Criteria: CNS metastases Active or suspected autoimmune disease 	 <u>Primary</u>: OS [Time Frame: 25 months] <u>Secondary</u>: ORR OS

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
A Master Protocol of Phase 1/2 Studies of Nivolumab in Advanced NSCLC Using Nivolumab as Maintenance After Induction Chemotherapy or as First-line Treatment Alone or in Combination with Standard of Care Therapies (CheckMate 370: Checkpoint Pathway and nivolumab Clinical Trial Evaluation 370) NCT02574078	RCT	April 2022	 Nivolumab Investigator's choice chemotherapy 	 N=1953 Age 18 years and older Inclusion Criteria: Stage IV NSCLC ECOG PS 0-2 Tumor tissue available for biomarker evaluation Exclusion Criteria: CNS metastases Suspected autoimmune diseases Hep B, Hep C, history of testing positive for HIV or AIDS 	 Primary: OS [Time Frame: 60 months] PFS [Time Frame: 48 months] Secondary: ORR DOR

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes	
Pembrolizumab						
A Randomized Open- Label Phase III Trial of MK-3475 Versus Platinum Based Chemotherapy in 1L Subjects with PD-L1 Strong Metastatic Non-Small Cell Lung Cancer NCT02142738	RCT	May 2018	 Pembrolizumab Cisplatin or carboplatin Gemcitabine or paclitaxel or pemetrexed 	 N=305 Age 18 years and older <u>Inclusion Criteria</u>: Stage IV NSCLC No driver mutations PD-L1 strong expression determined by IHC ECOG PS 0-1 Adequate organ function <u>Exclusion Criteria</u>: CNS metastases Autoimmune diseases 	 Primary: PFS [Time Frame: 2 years] Secondary: ORR OS 	

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
A Randomized, Open Label, Phase III Study of Overall Survival Comparing Pembroliz umab (MK-3475) Versus Platinum Based Chemotherapy in Treatment Naïve Subjects with PD-L1 Positive Advanced or Metastatic Non-Small Cell Lung Cancer (Keynote 042) NCT02220894	RCT	February 2018	 Pembrolizumab Carboplatin Paclitaxel or pemetrexed 	 N=1240 Age 18 years and older Inclusion Criteria: advanced or metastatic NSCLC No driver mutations PD-L1 positive determined by IHC ECOG PS 0-1 Adequate organ function No prior systemic chemotherapy Exclusion Criteria: CNS metastases Autoimmune diseases 	 <u>Primary</u>: OS [Time Frame: 2.5 years] <u>Secondary</u>: PFS
A Phase II/III Randomized Trial of Two Doses of MK- 3475 (SCH900475) Versus Docetaxel in Previously Treated Subjects with Non- Small Cell Lung Cancer NCT01905657	RCT	March 2019	 Pembrolizumab low/high dose Docetaxel 	 N=1034 Age 18 years and older <u>Inclusion Criteria</u>: NSCLC with progression after at least 2 cycles of platinum-containing doublets PD-L1 positive determined by IHC ECOG PS 0-1 Exclusion Criteria: prior docetaxel CNS metastases Active autoimmune disease History of HIV, Hep B or Hep C 	 <u>Primary</u>: OS, AEs [Time Frame: 3 years] <u>Secondary</u>: ORR DOR

Trial	Study Design	Estimated Completion	Comparators	Patient Population	Primary Outcomes
Atezolizumab					
A Phase III, Open- label, Multicenter, Randomized Study to Investigate the Efficacy and Safety of Atezolizumab Compared with Docetaxel in Patients with Non-Small Cell Lung Cancer After Platinum Failure [OAK] NCT02008227	RCT	June 2017	• Atezolizumab Docetaxel	 N=1225 Age 18 years and older <u>Inclusion Criteria</u>: Advanced or metastatic NSCLC progression during or after platinum-containing regimen measurable disease according to RECIST 1.1 ECOG PS 0-1 <u>Exclusion Criteria</u>: Prior docetaxel CNS metastases History of autoimmune disease 	 <u>Primary</u>: OS [Time Frame: 4.5 years] <u>Secondary</u>: ORR DOR PFS