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Chronological Trends in Progression-free, Overall and Post-progression Survival in First-line Therapy for Advanced Non-small-cell Lung Cancer

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1 **Chronological Trends in Progression-free, Overall and Post-progression**
2 **Survival in First-line Therapy for Advanced Non-small-cell Lung Cancer**

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4
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ABSTRACT

BACKGROUND: There is a debate about the merits of progression-free survival (PFS) versus overall survival (OS) as primary end points in non-small-cell lung cancer (NSCLC). It has been postulated that post-progression therapy may influence OS in both arms. To investigate this issue, we analyzed chronological trends in PFS and OS in advanced NSCLC using restricted mean survival times (RMSTs).

METHODS: We digitized survival curves from first-line phase III trials published between 1998 and 2015 in 13 leading journals to compute RMSTs for PFS and OS at three truncation landmarks (5, 12, and 18 months).

RESULTS: Among the 161 trials identified, RMSTs could be computed for both endpoints in 102, 97 and 82 trials for the 5, 12 and 18 months truncation landmarks, respectively. Post-progression survival (PPS) in the control arm, quantified as mean OS minus mean PFS truncated at 18 months, was on average 3.3 months between 1998 and 2003, 4.4 months between 2004 and 2009, and 5.4 months between 2010 and 2015. This increase was due to increasing RMST for OS over time, with no increase in RMST for PFS. The average within-trial difference in RMSTs between experimental and control arm was close to 0 for OS and less than 1 month for PFS.

CONCLUSIONS: There is a progressive increase in PPS in NSCLC trials, likely from salvage therapy. These results question both PFS and OS as sensitive end points in first-line trials, but suggest that the outlook for patients is improving regardless of within-trial gains.

48

INTRODUCTION

49 There has been significant progress in the treatment of non-small cell lung cancer
50 (NSCLC) from the identification and targeting of key molecular alterations—such as
51 epidermal growth factor receptor (EGFR) activating mutations, and ALK-EML4 and
52 ROS1 alterations¹⁻⁵—as well as from the use of immunotherapy.⁶⁻⁹ In parallel to
53 these advances, there has been a debate about the adequacy of overall survival
54 (OS) versus tumor-based end points —such as progression-free survival (PFS)—as
55 the most suitable and reliable primary end points for phase III trials in various tumor
56 types, including NSCLC.¹⁰ One of the issues raised in this debate is the possibility
57 that OS may be considerably influenced by treatments administered after disease
58 progression, and cross-over treatments in particular, which may dilute the treatment
59 effect.^{10, 11} Corroborating this concern, Hotta *et al*¹² and Hayashi *et al*¹³ have shown
60 that both median OS and post-progression survival (PPS) have increased along the
61 years, with absent or less pronounced simultaneous increases in median PFS in first-
62 line trials in advanced NSCLC. However, assessment of the relationship between
63 trends in PFS and OS over time are best made using mean survival times rather than
64 the medians, which do not have additive properties.¹⁴ In order to assess that
65 relationship, and in the attempt to quantify the magnitude of increase in PPS over the
66 years, we investigated trends in OS and PFS results during 18 years of clinical
67 research in NSCLC by using restricted mean survival times (RMSTs).

68

69

MATERIAL AND METHODS

70 Search strategy

71 We used PubMed, the medical subject headings ‘lung neoplasms’ and ‘drug therapy’,
72 and the filter option ‘randomized controlled trials’ to search for clinical trials on

73 systemic anticancer therapies published between January 1, 1998, and December
74 31, 2015 in 13 leading journals in the field of clinical oncology or lung cancer
75 specifically (*Annals of Oncology*, *British Journal of Cancer*, *Cancer*, *Chest*, *Clinical*
76 *Cancer Research*, *European Journal of Cancer*, *Journal of Clinical Oncology*, *Journal*
77 *of the National Cancer Institute*, *Journal of Thoracic Oncology*, *Lancet Oncology*,
78 *Lung Cancer*, *The Lancet* and *The New England Journal of Medicine*). We only
79 included phase III trials whose main results had been published in the study period.
80 We only included first-line treatment trials or those testing a maintenance strategy
81 after the first line. We excluded trials on supportive therapy alone, papers reporting
82 combined analyses of two or more separate trials already included in the study,
83 randomized phase II trials, and companion studies on correlative biology or
84 prognostic factors. Finally, we excluded randomized trials for which no phase was
85 explicitly stated if they had fewer than 100 evaluable patients per arm. With a cut-off
86 date of June 2016, we tried to obtain published OS results in later publications for the
87 trials that fell within the study period and for which OS results were immature when
88 the main paper was published.

89

90 **Data collection and definitions**

91 For each selected trial, we retrieved the main paper reporting the efficacy results and
92 abstracted the general characteristics of the trial and relevant data on OS and PFS,
93 time to tumor progression (TTP) or time to treatment failure (TTF). Despite their
94 slightly different meanings, we treated PFS, TTP and TTF together due to their
95 frequently overlapping definitions by investigators.¹⁵ When not explicitly stated, we
96 considered the primary end point as the metric used for sample-size calculation or
97 the end point first cited in the 'Methods' or 'Results' section of papers. We defined

98 trials with molecular selection of patients as those for which such selection was an
99 inclusion criterion.

100

101 **Statistical analysis**

102 The (unrestricted) mean survival (PFS or OS) of patients under a certain treatment
103 can be computed as the area under the Kaplan-Meier survival curve corresponding to
104 that treatment when the last patient under observation suffers the event of interest.

105 When the last patient is censored, the (unrestricted) mean survival cannot be
106 computed. In that case, the most frequently employed alternative is to compute the
107 RMST, which is the mean survival in the time interval between time 0 to a truncation
108 landmark, t . For the comparison of the RMST in two arms of a given trial, t is usually
109 chosen as the shortest time for which survival data are available for both arms. Since
110 our goal was to estimate PPS on the basis of (restricted) mean PFS and OS for a
111 large number of trials, we defined various arbitrary truncation landmarks to be used
112 across trials. For each trial, we identified the published Kaplan-Meier curves for PFS
113 (or TTP/TTF when this was the case) and OS, and digitized such curves using
114 CurveSnap.¹⁶ We used the same density of points (10/month) for all curves, thus
115 ensuring the same precision. Each curve was digitized in triplicate, once by each
116 different individual, and one of the authors (JR) selected manually the digitization that
117 appeared to better represent the curve shape. Using the digitized curves, we
118 computed RMSTs automatically, using software code created for the study purpose.
119 We chose three truncation landmarks (5, 12 and 18 months) based on the number of
120 curves that would provide sufficient information until such landmarks. We computed
121 RMST from the available PFS and OS curves from each trial and the average of
122 these RMSTs for each year of publication. The RMSTs thus computed indicate the

123 average (progression-free or overall) survival of patients in the control arm over the
124 period comprised by t . For the description of chronological trends, we used the
125 control arm in each trial for simplicity, under the assumption that improvements
126 obtained from clinical trials are reflected in the evolution of the control arms of future
127 trials. However, to provide a more complete view of chronological trends, we also
128 performed some analyses using the experimental arms of each trial. This was
129 straightforward in two-arm trials; for trials with more than two arms, we took the
130 average RMST (for PFS and OS) of the experimental arms. We quantified the
131 magnitude of mean PFS and OS gain as the difference in RMSTs between the
132 control and experimental arms. For the illustrative analysis involving median PFS and
133 OS times, we used the same set of trials for which RMSTs could be computed for
134 $t=12$ months, the landmark closer to the expected median OS in contemporary first-
135 line trials. In order to account for the potential influence of cross-over on OS, we
136 abstracted data from each trial about the presence of such cross-over in the control
137 and one experimental arm. We then conducted sensitivity analyses of trends in PFS
138 and OS over time according to the use of cross-over.

139

140

RESULTS

141 **General characteristics of the phase III trials**

142 The search yielded 161 phase III trials that were eligible for analysis. There were 144
143 trials testing a first-line therapy, and 17 in which the randomized intervention was
144 maintenance or treatment extension after first line. A total of 73,850 patients were
145 randomized to 353 treatment arms, with a median of 173 patients per arm. The yearly
146 mean number of patients per trial increased until 2009, when it started to plateau or
147 even decrease (Figure 1). This decrease was in part due to 11 trials with molecularly-

148 selected patients, all of which published after 2006: the median number of patients
149 per arm in these trials was 115, compared with 181 in trials with no molecular
150 selection. However, the decrease after 2009 may also have been due to increasing
151 use of PFS as primary end point, as this usually requires fewer patients (Figure 2).

152

153 The number of arms per trial was two in 137 cases, three in 17, and four in seven
154 trials (three of which with a factorial 2 x 2 design). Nineteen trials had a placebo-,
155 observation- or best supportive care-alone arm. Three trials had two co-primary end
156 points. In trials with a single primary end point, this was OS in 103 (64% of the total)
157 trials; PFS (N=26), TTP (N=1) or TTF (N=1) in 28 (17.4%) cases (henceforward
158 called PFS); response rate or clinical benefit rate in 16 (10.0%); quality of life or
159 toxicity in 10 trials (6.2%); and PFS without grade 4 toxicity in one (0.6%). Over the
160 18 years spanned by the study, the use of PFS as primary end point progressively
161 increased, with an accompanying decrease in the use of response or clinical benefit
162 rates (Figure 2).

163

164 **Time trends in mean and median OS and PFS times**

165 For $t=5$ months, computation of RMSTs was possible for 145 trials in the case of OS,
166 113 trials in the case of PFS, and 102 trials for both of these end points. For $t=12$
167 months, the RMST could be computed in 140 trials for OS and 107 trials for PFS;
168 computation of both RMSTs for the same trial was possible in 97 trials. Finally, for
169 $t=18$ months, the corresponding numbers of trials for which the RMSTs could be
170 computed were 122 for OS, 89 for PFS, and 82 for both end points. Table 1 shows a
171 descriptive comparison between trials for which RMSTs could or could not be
172 computed for PFS and OS truncated at 18 months. Trials for which RMSTs could be

173 computed were larger and published more recently, whereas the distribution of
174 primary end points was similar. The trials for which RMSTs could be computed were
175 also less likely to report gain in OS and more likely to report gain in PFS, but this may
176 be related to the fact that these trials were published more recently (and hence used
177 PFS as primary end point more frequently). Table 2 shows the number of trials for
178 each 2-year period of interest for which both of these means could be computed.

179
180 Figure 3 displays the evolution over time of PFS and OS in control arms, considering
181 the yearly average RMSTs with the 5, 12 and 18 months truncation landmarks; for
182 illustration, the average median times for PFS and OS are also shown. Not
183 surprisingly because of the short truncation landmark, there is little separation in the
184 curves that depict the yearly average mean PFS and OS for $t=5$ months (Panel A).
185 Such separation only becomes evident with the truncation landmarks of 12 (Panel B)
186 and 18 (Panel C) months. Moreover, a chronological trend for an increase in mean
187 OS with no accompanying increase in mean PFS becomes apparent with $t=12$
188 months, and more pronounced with $t=18$ months. When median times are
189 considered, a trend for increasing yearly average median OS with no accompanying
190 increase in PFS is also evident (Panel D). In this case, however, the degree of
191 separation may be misleading, because it does not directly indicate PPS.

192
193 Table 3 shows average RMSTs for PFS and OS in control arms for each 6-year
194 period spanned by the study, and according to the truncation landmarks of 5, 12 and
195 18 months. If the difference between RMSTs for OS and PFS over 18 months is used
196 to quantify PPS, the average duration of PPS was $8.9-5.6=3.3$ months in trials
197 published between 1998 and 2003, $10.0-5.6=4.4$ months in those published between

198 2004 and 2009, and $11.4-6.0=5.4$ months in those published between 2010 and 2015
199 (see also Figure 3, Panel C).

200

201 **Magnitude of mean PFS and OS gains over time**

202 Considering the trials for which RMSTs for both PFS and OS could be computed, the
203 average difference in RMSTs between experimental and control arms per 6-year
204 period and overall can be seen in Table 4. On average across trials, and regardless
205 of period or truncation landmark, there was essentially no gain in mean OS, whereas
206 the average gain in mean PFS was always less than 1 month. For $t=5$ months, the
207 differences in RMSTs between experimental and control arms within trials ranged
208 from a loss of 1 month to a gain of 2 months for PFS, and from a loss of 0.4 month to
209 a gain of 1.2 months for OS. For $t=12$ months, these differences ranged from a loss
210 of 1.8 months to a gain of 3.3 months for PFS, and from a loss of 1.2 months to a
211 gain of 1.9 months for OS. For $t=18$ months, these differences ranged from a loss of
212 1.9 months to a gain of 5.1 months for PFS, and from a loss of 1.6 months to a
213 maximum gain of 2.7 months for OS. When only trials with a significant gain in OS
214 are considered among those with sufficient data for analysis with $t=18$ months, the
215 average gain in OS was 1.2 month; in contrast, this average was 0 among trials with
216 no significant gain in OS.

217

218 **The potential influence of cross-over**

219 Cross-over was a feature of trial design or recommended by protocol in only six of
220 the 161 trials overall. All these trials had two arms and had the first-line therapy as
221 their main focus; in three of six cases, cross-over was recommended for both trial
222 arms, whereas in the other three cross-over was from the control to the experimental

223 arm (N=2) or vice-versa. When reported (N=4), the percentage of patients from the
224 control arm crossing-over to the same or similar agent as the experimental arm
225 ranged from 70.0% to 94.6%. For the remaining 155 trials, any post-progression
226 therapy was mentioned in 94 trials, 81 of which providing the percentages of patients
227 receiving such therapy. The median percentage of post-progression therapy was
228 42% in both control and experimental arms, with a clear increase in this percentage
229 over time (data not shown). Of the 94 trials in which post-progression therapy was
230 mentioned, cross-over could be characterized in 36 cases by looking at the
231 percentages of patients receiving the same or a similar agent as first-line therapy in
232 the post-progression setting. The median percentage of patients crossing-over was
233 18.5% in the control arms and 19.1% in the experimental arms. Of the total of 42
234 trials with cross-over (whether or not by design), 10 reported a significant gain in OS
235 (23.8%); among 119 trials without cross-over (or no mention thereof), a significant
236 gain in OS was reported in 27 (22.7%).

237
238 We assessed the evolution over time of yearly average RMST for PFS and OS in
239 control arms, considering the same 82 trials with sufficient data for the 18-month
240 truncation landmark, according to the presence of cross-over. We observed similar
241 trends for an increasing average OS and stable average PFS both in trials with no
242 reported cross-over (N=55) and in those with cross-over (whether or not by design;
243 N=27; data not shown). Among trials with cross-over, average PFS and OS according
244 to period were 5.7 months and 9.4 months for 1998 to 2003 (PPS=3.7 months); 5.3
245 months and 10.9 months for 2004 to 2009 (PPS=5.6 months); and 5.9 months and
246 12.6 months for 2010 to 2015 (PPS=6.7 months). Corresponding averages for trials
247 with no cross-over were 5.8 months and 9.0 months for 1998 to 2003 (PPS=3,2

248 months); 5.6 months and for 9.8 months 2004 to 2009 (PPS=4.2 months); and 6.1
249 months and 10.6 months for 2010 to 2015 (PPS=4.5 months).

250

251 Finally, we investigated trends in the average difference in RMSTs (with the 18-month
252 truncation landmark) between experimental and control arms within trials according
253 to the presence of cross-over. For trials with cross-over (N=27), the average
254 difference in RMSTs for PFS and OS were 0.3 month and 0.3 month for 1998 to
255 2003; 1.0 month and 0.4 month for 2004 to 2009; and 1.5 month and 0 month for
256 2010 to 2015. Corresponding differences for trials with no cross-over were 0.5 month
257 and 0 month for 1998 to 2003; 0.5 month and 0.2 month for 2004 to 2009; and 0.5
258 month and 0.2 month for 2010 to 2015. When the overall 18-year period is
259 considered, the average difference in RMSTs for PFS and OS were 1.1 month and
260 0.2 month, respectively, for trials with cross-over, and 0.5 month and 0.2 month,
261 respectively, for those without cross-over.

262

263

DISCUSSION

264 Our results disclose a progressive increase in mean OS with no obvious increase in
265 mean PFS following first-line therapy for advanced NSCLC over the years. Although
266 less appropriate from a methodological point of view, the same findings hold true
267 when these chronological trends are assessed using medians. Moreover, the
268 comparison of gains in mean PFS and OS times within trials show no substantial gain
269 on average. Therefore, our results collectively suggest that the outlook for patients
270 has clearly improved over the 18-year period of analysis, but not necessarily due to
271 improvements within trials. Rather, our data suggest that the increase in OS over

272 time is more likely due to post-progression therapy than to the treatments tested
273 within the trials.

274

275 In contrast with two previous smaller reviews that analyzed PFS and OS using
276 medians,^{12, 13} we ascertained trends in both end points using the RMST, a more
277 appropriate measure in the attempt to investigate differences in duration of OS and
278 PFS (*i.e.*, the duration of PPS),¹⁴ In a previous study, Trinquart *et al*,¹⁷ computed
279 RMSTs for the primary end point in 54 trials in several cancer types and therapeutic
280 settings, in an attempt to compare treatment effects estimated by the hazard ratio
281 (HR) and by differences and ratios of RMSTs. These authors pointed out that the HR
282 may overestimate the treatment effect as measured by the ratio of RMST. Several
283 other authors pointed to advantages of using RMSTs, including in settings where
284 hazards are non-proportional.^{14, 18, 19}

285

286 The extent to which our findings and conclusions are misleading due to the use of
287 RMSTs is unknown. One of the alternatives—the use of medians—is seen by many
288 as inadequate towards to goal of investigating PPS.¹⁴ Another alternative—the
289 computation of unrestricted means using observed data or parametric assumptions is
290 either not feasible or subject to the choice of models. These considerations
291 notwithstanding, confirmation of our findings would have important implications for
292 trial design in NSCLC, especially with regard to the choice of the primary end point.
293 There is an ongoing debate about which is the most adequate primary end point in
294 several tumor types, including NSCLC. We have argued that OS is generally
295 inadequate in settings for which there is salvage therapy,²⁰ which is increasingly the
296 case in NSCLC. The current results seem to confirm this contention, because there

297 has been essentially no gain in mean OS within trials, with a maximum gain of 2.7
298 months. On the other hand, we and others have argued in favor of PFS as primary
299 end point,^{10, 11} but the current results also question whether PFS is a sensitive end
300 point in this setting, because no substantial gain in mean PFS within trials could be
301 noted on average; in this case, however, the maximum mean gain in PFS was 5.1
302 months (for $t=18$ months). Once again, the extent to which the doubts about the
303 sensitivity of PFS and OS in our analysis are due to the use of RMSTs—as opposed
304 to conventional metrics—remains unknown. As shown in Table 1, significant gains in
305 PFS (as ascertained by conventional methods) were reported in 37% of trials,
306 whereas only 23% reported a significant gain in OS.

307
308 Interestingly, the current results show a progressive increase in OS despite
309 increasing use of PFS as primary end point (Figure 2) and no increased frequency of
310 OS gain usually ascertained by the logrank test (data not shown). Moreover, the
311 progressive increase in OS reported herein confirms previous analyses based on a
312 smaller number of trials and on real-life data.^{13, 21, 22} Finally, these results are in line
313 with the various recent instances in which a very active agent (as judged by
314 improvements in response rate or PFS) did not lead to improved OS, most likely
315 because of cross-over after disease progression.^{3, 4, 23}

316
317 Our study is subject to several limitations, the chief of which is the inability to
318 compute RMSTs for all eligible trials. Several authors have called for more extended
319 publication of results from phase III trials using RMSTs and other measures of
320 treatment effect that may complement the HR.^{14, 17, 18} It is hoped that increased
321 publication of such results will allow for more complete analyses in the future. A

322 second limitation is the potential for publication bias. Our analysis was restricted to
323 articles published in selected medical journals indexed in PubMed and within a
324 limited time period. This might have led to identification of phase III trials that are
325 systematically different from trials that have not been retrieved in our search.
326 However, we have consciously made this restriction, as the journals included in the
327 analysis report the vast majority of practice-influencing phase III trials. It is possible,
328 therefore, that we have analyzed studies with gains in OS preferentially. Another
329 limitation of our study relates to the inability to control for several variables that may
330 confound the associations and trends observed. For example, many recent trials
331 have included molecularly-selected patient populations with an improved prognosis;
332 the influence of these or even more subtle differences in patient profile on the current
333 results is uncertain. Such differences could possibly relate to tumor histology,
334 performance status, and other prognostic or predictive factors. Likewise, improved
335 trial methodology over the years may have influenced our results. Similarly, we
336 cannot ascertain the extent to which stage migration, other differences in patient
337 selection, or improvements in supportive care have influenced the apparent increase
338 in OS over time. Finally, incomplete reporting of information on cross-over and our
339 use of aggregated data—which precludes separate analyses of results among
340 patients with and without cross-over—prevent us from reliably ascertaining the
341 influence of cross-over on the trends for increasing OS without an accompanying
342 trend for increasing PFS. Nevertheless, our sensitivity analyses suggest that the
343 influence of cross-over is either not apparent or cannot be properly quantified, given
344 the similarity of results for (1) evolution over time of PFS and OS in control arms and
345 for (2) trends in average difference in RMSTs between experimental and control arms
346 within trials, when trials with and without cross-over are compared.

347

348 Our results may not be relevant for immunotherapy trials, only two of which were
349 represented in our sample.^{24, 25} In some of the randomized trials of antibodies against
350 the programmed death (ligand) 1 pathway, a significantly increased OS was not
351 always accompanied by an increased PFS.^{6, 9, 26} Since most immunotherapy trials
352 were published after our analysis period, our results apply essentially to
353 chemotherapy and targeted therapy. Despite these limitations, the current study is
354 the first to assess RMSTs in a large number of trials within one specific setting.

355

356 **Conclusions**

357

358 There is a progressive increase in PPS in NSCLC trials, but this appears to be more
359 likely a result of post-trial use of salvage therapy. These results question both PFS
360 and OS as sensitive end points in first-line trials, but suggest that the outlook for
361 patients is improving regardless of within-trial gains. It is hoped that population
362 enrichment and the use of targeted agents and immunotherapy with markedly
363 improved efficacy will allow for treatment effects that are larger than the ones
364 currently observed. If so, OS will continuously improve for patients inside and outside
365 of clinical trials.

366

367

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REFERENCES

- 375 1. Mok TS, Wu YL, Thongprasert S, et al. Gefitinib or carboplatin-paclitaxel in
376 pulmonary adenocarcinoma. *N Engl J Med* 2009;361:947-957.
- 377 2. Kwak EL, Bang YJ, Camidge DR, et al. Anaplastic lymphoma kinase inhibition in
378 non-small-cell lung cancer. *N Engl J Med* 2010;363:1693-1703.
- 379 3. Rosell R, Carcereny E, Gervais R, et al. Erlotinib versus standard chemotherapy as
380 first-line treatment for European patients with advanced EGFR mutation-positive non-small-
381 cell lung cancer (EURTAC): a multicentre, open-label, randomised phase 3 trial. *Lancet*
382 *Oncol* 2012;13:239-246.
- 383 4. Shaw AT, Kim DW, Nakagawa K, et al. Crizotinib versus chemotherapy in advanced
384 ALK-positive lung cancer. *N Engl J Med* 2013;368:2385-2394.
- 385 5. Sequist LV, Yang JC, Yamamoto N, et al. Phase III study of afatinib or cisplatin plus
386 pemetrexed in patients with metastatic lung adenocarcinoma with EGFR mutations. *J Clin*
387 *Oncol* 2013;31:3327-3334.
- 388 6. Borghaei H, Paz-Ares L, Horn L, et al. Nivolumab versus Docetaxel in Advanced
389 Nonsquamous Non-Small-Cell Lung Cancer. *N Engl J Med* 2015;373:1627-1639.
- 390 7. Brahmer J, Reckamp KL, Baas P, et al. Nivolumab versus Docetaxel in Advanced
391 Squamous-Cell Non-Small-Cell Lung Cancer. *N Engl J Med* 2015;373:123-135.
- 392 8. Reck M, Rodriguez-Abreu D, Robinson AG, et al. Pembrolizumab versus
393 Chemotherapy for PD-L1-Positive Non-Small-Cell Lung Cancer. *N Engl J Med* 2016.
- 394 9. Rittmeyer A, Barlesi F, Waterkamp D, et al. Atezolizumab versus docetaxel in patients
395 with previously treated non-small-cell lung cancer (OAK): a phase 3, open-label, multicentre
396 randomised controlled trial. *Lancet* 2017;389:255-265.

- 397 10. Soria JC, Massard C, Le Chevalier T. Should progression-free survival be the primary
398 measure of efficacy for advanced NSCLC therapy? *Ann Oncol* 2010;21:2324-2332.
- 399 11. Saad ED, Katz A, Hoff PM, et al. Progression-free survival as surrogate and as true
400 end point: insights from the breast and colorectal cancer literature. *Ann Oncol* 2010;21:7-12.
- 401 12. Hotta K, Kiura K, Fujiwara Y, et al. Role of survival post-progression in phase III
402 trials of systemic chemotherapy in advanced non-small-cell lung cancer: a systematic review.
403 *PLoS One* 2011;6:e26646.
- 404 13. Hayashi H, Okamoto I, Morita S, et al. Postprogression survival for first-line
405 chemotherapy of patients with advanced non-small-cell lung cancer. *Ann Oncol*
406 2012;23:1537-1541.
- 407 14. A'Hern RP. Restricted Mean Survival Time: An Obligatory End Point for Time-to-
408 Event Analysis in Cancer Trials? *J Clin Oncol* 2016;34:3474-3476.
- 409 15. Saad ED, Katz A. Progression-free survival and time to progression as primary end
410 points in advanced breast cancer: often used, sometimes loosely defined. *Ann Oncol*
411 2009;20:460-464.
- 412 16. CurveSnap. Available at <https://curvesnap.en.softonic.com/>.
- 413 17. Trinquart L, Jacot J, Conner SC, et al. Comparison of Treatment Effects Measured by
414 the Hazard Ratio and by the Ratio of Restricted Mean Survival Times in Oncology
415 Randomized Controlled Trials. *J Clin Oncol* 2016;34:1813-1819.
- 416 18. Royston P, Parmar MK. Restricted mean survival time: an alternative to the hazard
417 ratio for the design and analysis of randomized trials with a time-to-event outcome. *BMC*
418 *medical research methodology* 2013;13:152.
- 419 19. Uno H, Claggett B, Tian L, et al. Moving beyond the hazard ratio in quantifying the
420 between-group difference in survival analysis. *J Clin Oncol* 2014;32:2380-2385.

- 421 20. Saad ED, Buyse M. Statistical controversies in clinical research: end points other than
422 overall survival are vital for regulatory approval of anticancer agents. *Ann Oncol*
423 2016;27:373-378.
- 424 21. Morgensztern D, Waqar S, Subramanian J, et al. Improving survival for stage IV non-
425 small cell lung cancer: a surveillance, epidemiology, and end results survey from 1990 to
426 2005. *J Thorac Oncol* 2009;4:1524-1529.
- 427 22. Breathnach OS, Freidlin B, Conley B, et al. Twenty-two years of phase III trials for
428 patients with advanced non-small-cell lung cancer: sobering results. *J Clin Oncol*
429 2001;19:1734-1742.
- 430 23. Fukuoka M, Wu YL, Thongprasert S, et al. Biomarker analyses and final overall
431 survival results from a phase III, randomized, open-label, first-line study of gefitinib versus
432 carboplatin/paclitaxel in clinically selected patients with advanced non-small-cell lung cancer
433 in Asia (IPASS). *J Clin Oncol* 2011;29:2866-2874.
- 434 24. Alfonso S, Valdes-Zayas A, Santiesteban ER, et al. A randomized, multicenter,
435 placebo-controlled clinical trial of racotumomab-alum vaccine as switch maintenance therapy
436 in advanced non-small cell lung cancer patients. *Clin Cancer Res* 2014;20:3660-3671.
- 437 25. Giaccone G, Bazhenova LA, Nemunaitis J, et al. A phase III study of
438 belagenpumatucel-L, an allogeneic tumour cell vaccine, as maintenance therapy for non-small
439 cell lung cancer. *Eur J Cancer* 2015;51:2321-2329.
- 440 26. Herbst RS, Baas P, Kim DW, et al. Pembrolizumab versus docetaxel for previously
441 treated, PD-L1-positive, advanced non-small-cell lung cancer (KEYNOTE-010): a
442 randomised controlled trial. *Lancet* 2016;387:1540-1550.
- 443

Table 1. Characteristics of trials with and without available RMSTs truncated at 18 months.

Characteristic	Trials with available RMSTs (N=82)	Trials without available RMSTs (N=79)
Year or publication		
1998 to 2003	19 (23%)	28 (35%)
2004 to 2009	27 (33%)	32 (41%)
2010 to 2015	36 (44%)	19 (24%)
Patients per arm, median (IQR)	183 (138; 304)	154 (99; 222)
Primary end point*		
Overall survival	55 (69%)	48 (61%)
Progression-free survival	14 (18%)	14 (18%)
Other	11 (14%)	16 (20%)
Reported gain in overall survival**	16 (20%)	21 (27%)
Reported gain in progression-free survival**	33 (40%)	20 (32%)

Percentages are rounded to integers. IQR, interquartile range; RMST, restricted mean survival time.

*Trials with co-primary endpoints not shown. **Denominator is the number of trials with information on gains, which we deemed to be present if there was a statistically significant difference in any pairwise comparison (or factor comparison in factorial trials) for PFS or OS following the trialists' definitions, regardless of which of these was the primary end point and of the number of arms in a given trial, and regardless of whether the difference favored the experimental or the control arm.

Table 2. Availability of data on mean PFS and OS for each 2-year period.

Year of publication	Number of trials	Number of trials with data for analysis according to truncation time (t)		
		For $t=5$ months	For $t=12$ months	For $t=18$ months
1998-99	12	6 (50%)	5 (42%)	4 (33%)
2000-01	15	6 (40%)	4 (27%)	3 (20%)
2002-03	20	13 (65%)	13 (65%)	12 (60%)
2004-5	22	16 (73%)	16 (73%)	12 (55%)
2006-07	17	9 (53%)	8 (47%)	6 (35%)
2008-09	20	10 (50%)	10 (50%)	9 (45%)
2010-11	19	14 (74%)	14 (74%)	13 (68%)
2012-13	21	17 (81%)	16 (76%)	15 (71%)
2014-15	15	11 (73%)	11 (73%)	8 (53%)
Total	161	102 (63%)	97 (60%)	82 (51%)

Table 3. Average RMSTs (in months) for each 6-year period of interest, control arms.

Years of publication	Average according to end point and truncation time (t)					
	For $t=5$ months (N=102)		For $t=12$ months (N=97)		For $t=18$ months (N=82)	
	PFS	OS	PFS	OS	PFS	OS
1998 to 2003	3.4	4.2	5.1	7.7	5.6	8.9
2004 to 2009	3.6	4.4	5.1	8.3	5.6	10.0
2010 to 2015	3.7	4.5	5.4	9.0	6.0	11.4

N indicates number of trials with available data.

Table 4. Average difference in RMSTs (in months) between experimental and control arms within trials.

Years of publication	Average difference according to end point and truncation time (t)					
	For $t=5$ months (N=102)		For $t=12$ months (N=97)		For $t=18$ months (N=82)	
	PFS	OS	PFS	OS	PFS	OS
1998 to 2003	0.3	0.1	0.3	0	0.3	0.1
2004 to 2009	0.1	0	0.5	0.1	0.7	0.2
2010 to 2015	0.2	0	0.7	0.1	0.9	0.1
Overall in 18 years	0.2	0	0.5	0.1	0.7	0.2

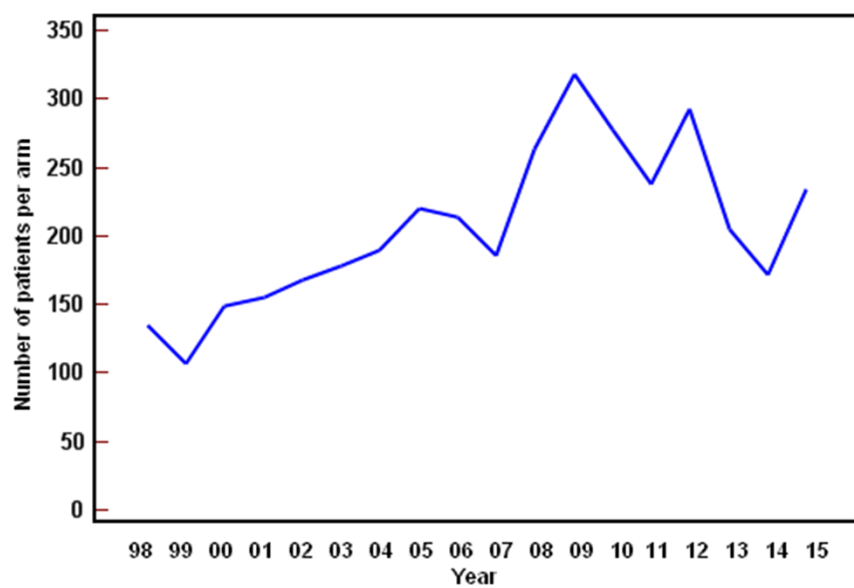
N indicates number of trials with available data.

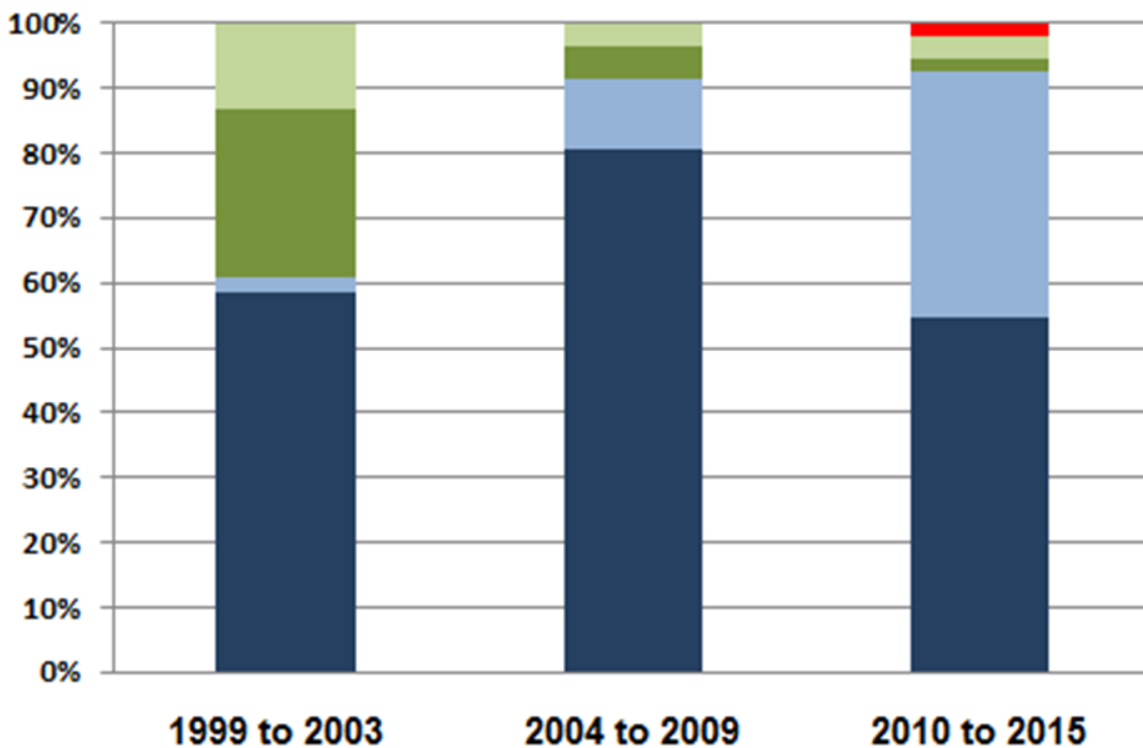
FIGURE LEGENDS

Figure 1. Yearly mean number of patients per arm over time.

Figure 2. Frequency of use of overall survival (dark blue), progression-free survival (light blue), response or clinical benefit rates (dark green), quality of life or toxicity (light green), and other primary end points (red) in the three 6-year periods spanned by the study.

Figure 3. Time trends in restricted mean and in median progression-free and overall survival times. Panel A, yearly average of the restricted mean survival times (RMSTs) for progression-free (PFS) and overall survival (OS) in control arms, truncated at 5 months; Panel B, yearly average of the RMSTs for PFS and OS in control arms, truncated at 12 months; Panel C, yearly average of the RMSTs for PFS and OS in control arms, truncated at 18 months; Panel D, yearly average of the median PFS and OS in control arms.





ACCEPTED M

