Statistical considerations and endpoints for clinical lung cancer studies: can progression free survival (PFS) substitute overall survival (OS) as a valid endpoint in clinical trials for advanced nonsmall-cell lung cancer?

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> Abstract: In the last decades significant progress has been achieved in the biological understanding of nonsmall-cell lung cancer (NSCLC) and its tumor heterogeneity has become more evident. The identification of novel tumor targets with different pathways has stimulated the search for anti-tumor agents with a specific target directed mode of action, stipulating the need of testing these agents in clinical trials with an appropriate choice of the study endpoint. Gold standard as an endpoint has been so far overall survival (OS). By definition there are 3 categories of classical endpoints applied generally in clinical lung cancer studies: survival time endpoints, symptom endpoints, and endpoints relying on patients' reporting. Beside classical endpoints like OS which are tending to show the direct clinical effect of treatment, efforts have been taken to substitute these classical endpoints by surrogates. As a surrogate candidate for OS progression-free survival (PFS) should have the inherent considerable advantage, that it can detect subpopulations with longer PFS intervals early. Based on the (sub-) population treated and having in mind the risk-benefit profile of the drug under consideration, PFS can be considered for regulatory decision making. If accompanied by some independent measures like quality of life or treatment toxicity, PFS should be able to cover the clinical benefit achieved by treatment. Selecting PFS as primary endpoint in Phase III trials of advanced NSCLC may be based on a number of questions such as: Does the definition of PFS fit into the setting used by other trials? Are there accepted consensus standards? Are there consistent surveillance intervals? Is validation for each agent group planned? Is the incremental improvement of PFS big enough (\geq 30%)? And are there some additional measures to confine clinical benefit? OS is still accepted as the gold standard in trials investigating advanced NSCLC. OS is easy to measure and precise but it may be difficult to interpret if treatment action takes place only in a small subinterval of overall survival. PFS with some additional measures has become attractive when it seems advisable to make study results available earlier. Candidates for supporting PFS as "additional measures" may be treatment toxicity and quality of life measures. PFS allows a more precise detection and attribution to effects of the investigational treatment without being compromised by subsequent treatments. Therefore "enriched PFS" can be considered as an alternative primary endpoint replacing OS in studies investigating advanced NSCLC. The endpoint selection process should always be performed carefully considering all true and surrogate endpoint options in respect to the hypotheses to be proven.

Keywords: Non-small-cell lung cancer; clinical trial; study endpoint; overall survival; progression-free survival



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Introduction

Non-small-cell lung cancer (NSCLC) is a wide spread disease with a proportion of about 80% in all 1.2 million lung cancers cases per year and is one of the leading causes of malignancy related to morbidity and mortality in the industrialized countries with increasing incidence worldwide (1). In most patients lung cancer is diagnosed in an advanced state with poor prognosis. Because in the last decade's significant progress has been achieved in the biological understanding, tumor heterogeneity of NSCLC became more evident and calls for treatment individualization (2). The identification of novel tumor targets with different pathways has stimulated the search for anti-tumor agents with a specific target directed mode of action. Taking into account subpopulations of patients and special efficacy pathways of new drugs, the gold standard overall survival as an adequate endpoint for clinical trials may have become questionable and has been challenged lately by endpoints not dependent on the growing number of subsequent therapies (3). Therefore, we will discuss in this paper the role of progression free survival as a possible alternative endpoint to overall survival in treatment of NSCLC (4-7).

Methods

Classical study endpoints

Based on the practice in the last centuries there are by definition three categories of classical endpoints generally applied in clinical studies: Survival time endpoints, symptom endpoints, and endpoints relying on patients' reporting. Adjacent to classical endpoints which are measuring single parameters and are tending to show the direct clinical effect of treatment, efforts have been taken to substitute these classical endpoints by surrogates (8-10). Motivated is the use of surrogate endpoints mainly by faster approval processes, smaller sample sizes, positive cost effects and shorter trial duration. The process of substituting a classical endpoint means to replace a rare endpoint or endpoint, describing a late event in the course of disease by a more frequent endpoint or endpoint describing an early event (11,12). It is widely accepted that such an endpoint should be ideally based on a biological rationale and should be related to the study endpoint of interest (13). It may be convenient for clinical trials to base the main hypothesis on tumor parameters, treatment toxicity, and drug pharmacology. The ultimate aim in the majority of patients however, is to prolong the remaining lifespan and reduce tumor morbidity without worsening the performance of the patient by treatment related adverse events, what is commonly understood as clinical benefit.

Survival time endpoints

Lifespan intervals such as progression-free survival (PFS) and

overall survival (OS) are the most common endpoints especially in Phase III studies (14,15). OS measures the clinical benefit of a treatment in total and is used for the regular approval. It is defined as the lifespan starting from recruitment (or first diagnosis of treated disease, or start of first treatment under consideration) to the event of death, what reason so ever. In a study with more than one treatment arm randomization of the study is essential (16). Blinding of the study is not necessary, especially in lung cancer treatments, since side effects are most known and co-medication is described. In ad-on treatment with biologics blinding may be possible (17,18). Censored are patients alive or lost to follow up (19). PFS has the same starting points as OS and includes in its endpoint measurement all deaths, whatever the reason is and all progressions of the tumor under investigation as well as new tumor manifestations. Censored are the patients without such events and those in the lost to follow up situation (20,21). Uncommon survival time endpoints include time to progression (TTP), disease free survival (DFS), recurrence free survival (RFS), and time to treatment failure (TTF) which will not be discussed in the sequel.

Tumor symptom endpoints

Leading symptoms that are directly related to the tumor mass and the extension of the disease can be used as a measure of treatment effect. In Phase III trials they are not so often employed as survival endpoints and defined by rates in a given time interval. The objective response rate (ORR) is a counting measure for the best treatment result (22). The occurrence of partial or complete remission of the tumor is counted as a success and the duration of remission is not taken into account. The number of objective tumor remissions is measured in relation to all evaluable patients in the study at a given time point (23,24). The change in tumor related symptom endpoints such as pain, weight loss, performance status, and dispnoea can measure the clinical benefit. Studies using these endpoints should be randomized blinded trials. Here, the measurement concerns the perspective of direct clinical benefit for each patient and can also measure the toxic side effects of the study medication (25). Blinding is often difficult (skin rash from the investigated drug!) and data are frequently missing (less evident events) or arrive incomplete (because of the amount of data) in the study center (18).

Patient reported outcomes

Simplicity, instantaneous availability, and proximity to the true health condition of patients are the main advantages in using patients' self-evaluation of their health performance in structured reports for clinical outcome description (26). Because linking patients' information directly to clinical benefit, quality of life (QOL) is important (27). In Europe the questionnaire EORTC

Table 1 Drugs and approval endpoints (33)

Drugs/regimens and approval endpoints	
First-line NSCLC	
Vinorelbine monotherapy	OS, ORR
Vinorelbine + cisplatin	OS, ORR
Docetaxel + cisplatin	OS, TTP, ORR
Gemcitabine + cisplatin	OS, TTP, ORR
Bevacizumab [†] + paclitaxel/carboplatin	OS
Paclitaxel + cisplatin	TTP, ORR, OS
Pemetrexed in combination with cisplatin ^{t,‡}	OS
Pemetrexed as maintenance therapy †	OS
Second-line NSCLC	
Docetaxel	OS, TTP, ORR
Erlotinib	OS, TTP, ORR
Pemetrexed ^{†,‡}	Durable ORR, decreased toxicity
Third-line NSCLC	,
Erlotinib	OS, TTP, ORR
Gefitinib [‡]	Durable ORR
ter or en	

[†]Limited to non-squamous non-small cell lung cancer; [‡]Accelerated approval; OS, overall survival; ORR, objective response rate; TTP, time to progression; NSCLC, non-smallcell lung cancer.

QLQ C30 is a widely used instrument translated into 81 languages and employed in more than 3,000 clinical trials (28). The main questionnaire is supplemented by disease specific modules, so LC13 for lung cancer, where 13 special questions concerning lung cancer symptoms are assembled (29,30). In the United States a lung cancer symptom scale (LCSS) and a module of the American Thoracic Society are in application. LCSS was designed as a sitespecific measure of QOL for particular use in clinical lung cancer trials evaluating six major tumor symptoms (31). The questionnaire FACT-G (Functional Assessment of Cancer Therapy) is distributed by the American Thoracic Society and includes the special lung cancer module FACT-L (32).

Results

Application of endpoints in clinical studies

One of the main sources worldwide for a standardized procedure of regular drug approval is the FDA. In the FDA guidance draft of June 2011 concerning recommendations for clinical trial endpoints for the approval of anticancer agents three commonly used efficacy endpoints in assessing anti-neoplastic agents are advocated in

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NSCLC treatment: OS, PFS, and ORR (19). Nevertheless, the majority of approvals rely on OS. Of 16 trials which have been conducted for drug approval, 11 were based on OS as leading endpoint, 2 on PFS, 2 on durable ORR, and 1 on improvement in disease related symptoms, respectively. It was recently stated that the regular basis of approval for drug in treatment of advanced NSCLC has been OS (Table 1) (33). In the FDA perspective, studies have demonstrated that PFS as well as ORR may not reliably predict for a corresponding effect on survival (34,35). Not every detail is clear how to assess a risk/benefit profile that was stipulated for approving new drugs (36). However, it was claimed that an improvement in OS in randomized controlled trials has been the standard for establishing clinical benefit for drug approvals in advanced NSCLC. Not discussed in detail by FDA is the influence of subsequent treatments and special supportive care on OS. It is still hoped that the law of large numbers will rule out differences in comparisons of treatment arms in randomized trials.

Recent advances in molecular biology have led to the development of many anticancer agents targeting on specific target aberrant pathways and proteins of tumor cells. The number of agents available for testing asks for a more efficient and quicker drug testing system (12,37). Surrogate endpoints, which can replace or supplement traditional endpoints, are useful if they can be measured earlier, more conveniently or more frequently than true endpoints (38).

FDA allows accelerated approval on less-established surrogate endpoints, that are reasonably likely to predict clinical benefit and it is explicitly stated that this approval mechanism may only be used when the drug provides a benefit over available therapy (7). If a variable in a clinical study is correlated to a true clinical outcome that does not mean, that it can be used as a valid surrogate endpoint and replace the true clinical outcome (37). FDA still believes that QOL measures can be important clinical benefit endpoints, particularly in a predominantly symptomatic disease such as NSCLC (19). In these patients, with limited survival expectations, symptom palliation, quality of life, and convenience of therapy are especially important endpoints (39). The main argument for these instruments is that health care professionals are not accurate in evaluating subjective or palliative benefits associated with anticancer treatments, when compared with patient self-reports. But so far no approval has been based on QOL only. Contradictive facts in using QOL are blinding the trial, reliability and validity of instruments, and the handling of missing data. Since for OS the distinction between two or more treatments will require a larger study sample to show statistically significant differences, PFS is increasingly used as endpoint for randomized Phase II/III studies especially for testing schedules and regimens of targeted therapies.

PFS assessment of disease progression is based on radiologic testing at scheduled time points in cycles and the task is, to deliver reliable results for decision. In statistical models of survival



Figure 1 Methods of data handling with missing data on progression.



Figure 2 All events of OS are covered by PFS; PFS shows additionally the events of tumor control.

analysis as PFS and OS, not for all subjects the time-to-event will be observed, which will then be censored in the describing statistics. Censoring can be performed in several ways and in *Figure 1* different handling methods are demonstrated (20,40). The graphical output of survival data are best presented by Kaplan-Meier (KM) curves where the percentage of survived individuals is plotted against time, which is more reliable than individual survival probabilities. Reliability of KM curve diminishes with increasing time. But there is generally no defined time point or number where the tail of the curve becomes uncertain in interpretation.

At least nine events are related in assessing PFS: (I) Local regional relapse; (II) Distant metastases; (III) Second primary tumor-same entity; (IV) Second primary tumor-other entity; (V) *Death related to tumor; (VI) *Death related to another tumor; (VII) *Tumor independent death; (VIII) *Death related to tumor therapy; (IX) Lost to Follow-up: Censored. All events marked by * are also covered by OS (Figure 2) (41).

To demonstrate the impact of sample size *Figure 3* illustrates calculation for trials where the surplus in overall survival varies. Sample size is calculated by the log rank test (Freedman) under the assumption that in the two groups the hazards are proportional (PASS 11, version 11.0.7, 2011) (42,43). Further on, it is assumed that the survival times are exponentially distributed. If the median survival of the control group is 12 months then the 1-year survival rate is 50%. An increase of 10% to a 1-year rate of 55% is equivalent to a survival of 14 months. A rate of 60% corresponds to 16.3 months, 65% to 19.4 months, and 70% to 23.4 months, nearly doubling the survival time. For superiority the one-sided test is used. The power was set to 90% or 80% and α =0.025 or α =0.05. Accrual is 1:1 for the two groups. The sample sizes vary between N=4059 and N=151 depending on the power, significance and survival proportion. A small incremental increase in survival



Figure 3 Sample size calculation for proportion of 1-year survival of 0.55-0.70 of experimental group versus 1-year survival of 0.5 for controls in a 1:1 recruitment and with power of 80% or 90% and α =0.025 or α =0.05.

time accompanied by small error rates (power and $\boldsymbol{\alpha})$ leads to a large study population.

Substitution of endpoints

The main factor for PFS to predict OS is the duration of PFS (43). A significant survival gain can be predicted in a trial of 750 patients if 18% of patients responding to treatment. 500 patients are needed if 21% of patients respond and if it is postulated that 30% patients respond, at least 250 patients must be enrolled.

The necessary incremental gain for PFS to predict a survival improvement was a median of 1.7 months for trials with 750 patients, 2.2 months for 500 patients, and 3.3 months for trials with 250 patients. Hence, a significant surplus on PFS can only be predictive for OS if the sample is big enough. Statistically it comes to compare PFS and OS as two failure time distributions in the presence of censoring, where the distribution is shifted due to the different time interval of measurement (44). Compared to OS in PFS more competing risks are included (*Figure 2*). PFS as surrogate endpoint for OS can also be investigated by the aid of a stochastic model (45). The Process was demonstrated theoretically by a statistical OS model with two independent stochastic processes defining an interim and a restart point of progression. Since it has been discovered that the interval after progression to death may have similar distribution for most of the lung cancer trials PFS as a surrogate of OS seem to be reasonable (46,47). The strongest prediction of PFS for OS results from a high correlation of PFS and OS and a high treatment effect homogeneity (*Figure 4*).

Quality of life assessment represents an independent outcome in clinical research and an index of effectiveness of a treatment (48,49). The augmentation of PFS by QOL is a combination of two almost independent endpoints depicting a patient selfevaluation of personal health performance and an objective management. Subsequent chemotherapy improvement of tumor related symptoms translates to a better quality of life. Using QOL questionnaires can result in well documented health-related quality of life data and complementary efficacy information, helping in clinical decision finding (50).

Endpoints for targeted drugs

Based on the fact that conventional treatment of NSCLC has reached a plateau of effectiveness in improving the survival new targeted drugs have been developed. Testing targeted therapeutic agents in clinical trials need specific consideration for a number of reasons (51,52). The mode of action of the targeted agent depends specifically on molecular biology and therefore needs an adaption of the study design according to the tumor target concerning study endpoints, patient selection, diagnostic requirements, time frame, etc (53,54). The large number of targeted agents available for



Figure 4 OS prediction depends on the treatment effect homogeneity and the correlation with PFS.

clinical testing certainly is a challenging task in clinical research, which means also to solve specific methodological, logistic, administrative, and statistical issues (55).

If a targeted agent is tested against a placebo arm, and considering a longer time interval for observing progression events, it has been recognized that the drug development is time consuming even using PFS. An outcome-based adaptive randomization trial design for patients can be employed with conjunction of an early stopping rule and can be used to identify effective agents and match those treatments with patients' biomarker profiles (56). Testing the target may cause additional costs and logistic strength (e.g., EGFR-mutation testing). Additionally, to the limitations of PFS as the main endpoint are the limitations of the diagnostic test procedure itself that may lead to misinterpretation of study outcome and in the last consequence to an un-appropriate therapy (40).

Discussion

Merits and shortcomings of study endpoints

So far, OS was used as a gold standard in Phase III lung cancer studies but that concept may not work for the approval of chemotherapeutic drugs and targeted agents. For the choice of OS as endpoint there are two main pitfalls: I. the competition of death events others than caused by the treated lung cancer disease and the difficulty to distinguish between cancer related and not related deaths and II. the probable long time interval after end of treatment and death of the patient where many lifespan influencing events can happen (57). If the sample size is big enough the statistical rule of the "law of large number" will balance competing risks and under these assumptions overall survival can be accepted as a universal primary endpoint in lung cancer even if the potential causes of other deaths cannot be neglected. More complicated is the statistical handling of a possible long time interval after the end of treatment till death, with the determinants therapy effectiveness, subsequent therapies and salvage therapies making the process not any longer strictly depend directly on the primary treatment action taken (58). Hence, the aspect of respond to primary treatment is not depicted generally in OS. A numerical evaluation of subsequent treatments and influence correcting statistical analysis is almost not possible and depends on the tumor type. Since subsequent treatments are very divers these influences cannot be easily categorized (59-61).

Clearly, progression under treatment is a sign that the disease is not positively affected by treatment action and states the failure of treatment (62). If curative treatment is not available the absence of progression may be evaluated as partial success especially in case of mild side effects of the underlying treatment. For the patient itself non-progression may be an important signal that the disease has come to a stop with all psychological positive side effects. PFS is the clinical measure of this process.

PFS as measure of response or shrinkage of the tumor can act as an indicator of overall survival. But interpretation remains difficult for various reasons: Many patients showing no progression and no tumor size reduction to treatment, midterm side effects disturb the enduring effect of response and diminishing a beneficial overall survival, and no adequate further treatment options may be available in case of progression (63-65). Measurement of PFS may be constricted by errors in assessing progression and the fixed time schedule of reassessment of the tumor (65). In addition, there is no standardization of measurement in the chosen time interval, tumor type, and tumor mass. As a matter of fact, most of the progressions will happen between the therapy assessments or follow up intervals (66).

PFS as an endpoint is a choice worth considering if multiple therapy options exist. Then it can be compared if only the response of the study drug is measured (67). But PFS is not statistically validated as a surrogate for survival in all settings (68). It can happen, that the results of PFS on the one side and OS on the other side are contradicting. The measurement of progression is not as precise as statement of death in OS and may be subject of assessment bias particularly in open-label studies. Additionally, the definitions in various studies may vary and so it will be complicated to compare trials with different PFS definitions. The diagnostic complexity and costs are high, frequent radiological or other assessments are necessary. Important is the balanced timing of assessment within the study and especially between treatment arms. Sample size is often smaller in PFS in comparison with OS. This fact together with earlier results is an advantage for PFS. But the time interval between progression and death might be large and influence the ability of translating PFS into OS.

Endpoints for lung cancer trials

As in any other cancers, lung cancer trials must determine the clinical benefit of a specific new treatment. The concept of survival time-not necessarily OS-can be used for hypothesis driven statistics as a primary study endpoint and for the sample size calculation. The main thought of these advisements is that all positive actions taken, will translate into a longer overall survival. In the case that the study drug was given only for a short time period all other effects will be ruled out by the law of large numbers, in comparison with a similar population without investigational treatment (control arm). These thoughts are independently true whatever survival time concept is used (69,70).

Following the concept of evidence based medicine the clinical study should have a reasonable hypothesis, which incorporates a primary endpoint to be able to demonstrate the difference between the investigational and the standard treatment at a statistical significant level. The final evidence is simply provided by testing the given hypothesis. The choice of the endpoint definitely influences the treatment and study success (71). As a basic claim, success should always be postulated as the final result of the study. The chosen endpoint is a determinant of the sample size and has to be selected prospectively in an explicit manner and cannot be altered during the course of the trial. In advanced lung cancer the horizon of the remaining lifespan is quite limited. Thus it is not inequitable that PFS derived under the study drug will translate directly into OS (72).

Lung cancer can be taken as a chronic disease diagnosed often in an advanced stage with a remaining median lifespan of 7 to 15 months. Thus, in advanced lung cancer OS should be a quite natural measure of treatment success (73). Tumor progression is an indicator for disease worsening, a time point for new treatment considerations, and a negative signal for tumor morbidity and death. PFS is measured in the complete study population without exceptions and is an indirect measure for treatment benefit. The main advantages of PFS over OS are that the outcome measure is available earlier, that it can shorten the drug development time, and that there is no influence by subsequent therapies.

PFS as a surrogate for OS should have the inherent considerable advantage, that it can detect subpopulations with longer PFS intervals early. Based on the (sub-) population treated and having in mind the risk-benefit profile of the drug under consideration, PFS can be considered for regulatory decision making. If accompanied by some independent measures like QOL or treatment toxicity, PFS should be able to cover the clinical benefit achieved by treatment. In a recent report on surrogate endpoints in oncology by the German Institut für Qualität und Wirtschaftlichkeit im Gesundheitswesen (IQWiG) two levels of evaluation were communicated: The validity of the surrogate endpoint should be evaluated and the conclusions should be drawn from the seen effects (74). Selecting PFS as primary endpoint in Phase III trials of advanced NSCLC may be based on a number of questions such as: Does the definition of PFS fit into the setting used by other trials? Are there accepted consensus standards? Are there consistent surveillance intervals? Is validation for each agent group planned? Is the incremental improvement of PFS big enough (\geq 30%)? And are there some additional measures to confine clinical benefit?

Conclusions

OS as a concept of clinical benefit is still accepted as the gold standard in trials investigating advanced NSCLC. OS is easy to measure and precise but it may be difficult to interpret if treatment action takes place at the beginning of the measured survival interval. PFS, basically, does not overcome those difficulties of OS. However, PFS with some additional measures has

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become attractive when it seems advisable to make study results available earlier. Candidates for supporting PFS as "additional measures" may be treatment toxicity and quality of life measures. PFS allows a more precise detection and attribution to effects of the investigational treatment without being compromised by subsequent treatments. Therefore "enriched PFS" can be considered as an alternative primary endpoint replacing OS in studies investigating advanced NSCLC. In comparison to OS, PFS is clearly more complex to assess, requires a very stringent definition in clinical trials and as a consequence the ability for a more intensive radiological imaging. Nevertheless, the process of endpoint selection should always be performed carefully judging all true and surrogate endpoint options in respect to the study hypotheses including the projected study design.

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