Guidance for Industry Clinical Trial Endpoints for the Approval of Cancer Drugs and Biologics

DRAFT GUIDANCE

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> U.S. Department of Health and Human Services Food and Drug Administration Center for Drug Evaluation and Research (CDER) Center for Biologics Evaluation and Research (CBER)

> > April 2005 Clinical/Medical

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Guidance for Industry¹ Clinical Trial Endpoints for the Approval of Cancer Drugs and Biologics

This draft guidance, when finalized, will represent the Food and Drug Administration's (FDA's) current

thinking on this topic. It does not create or confer any rights for or on any person and does not operate to

bind FDA or the public. You can use an alternative approach if the approach satisfies the requirements of

the applicable statutes and regulations. If you want to discuss an alternative approach, contact the FDA

staff responsible for implementing this guidance. If you cannot identify the appropriate FDA staff, call

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19 I. INTRODUCTION20

21 This guidance provides recommendations to sponsors on endpoints for cancer clinical trials

submitted to the FDA to support effectiveness claims in new drug applications (NDAs),

biologics license applications (BLAs), or supplemental applications.²
 24

the appropriate number listed on the title page of this guidance.

The FDA is developing guidance on oncology endpoints through a process that includes public
workshops of oncology experts and discussions before the FDA's Oncologic Drugs Advisory
Committee (ODAC).³ This guidance is the first in a planned series of cancer endpoint
guidances. It provides background information and discusses general regulatory principles.

29 Each subsequent guidance document will focus on endpoints for specific cancer types (e.g., lung

30 cancer, colon cancer) to support drug approval or labeling claims. The endpoints discussed in

31 this guidance document are for drugs to treat patients with an existing cancer. This guidance

32 does not address endpoints for drugs to prevent or decrease the incidence of cancer.

33

34 FDA's guidance documents, including this guidance, do not establish legally enforceable

35 responsibilities. Instead, guidances describe the Agency's current thinking on a topic and should

36 be viewed only as recommendations, unless specific regulatory or statutory requirements are

¹ This guidance has been prepared by the Division of Oncology Drug Products and the Division of Therapeutic Biologic Oncology Drug Products in the Center for Drug Evaluation and Research (CDER) in cooperation with the Center for Biologics Evaluation and Research (CBER) at the Food and Drug Administration.

 $^{^{2}}$ For the purposes of this guidance, all references to *drugs* include both human drugs and biological products unless otherwise specified.

³ Transcripts are available at http://www.fda.gov/cder/drug/cancer_endpoints/default.htm.

cited. The use of the word *should* in Agency guidances means that something is suggested orrecommended, but not required.

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II. BACKGROUND

41 42

43 Clinical trial endpoints serve different purposes. In conventional oncology drug development, 44 early phase clinical trials evaluate safety and identify evidence of biological drug activity, such 45 as tumor shrinkage. Endpoints for later phase efficacy studies evaluate whether a drug provides 46 a clinical benefit such as prolongation of survival or an improvement in symptoms. The 47 following sections discuss the general regulatory requirements for efficacy and how they have 48 influenced endpoint selection for the approval of cancer drugs. Later sections describe these 49 endpoints in more detail and discuss whether they might serve as measures of disease activity or 50 clinical benefit in various clinical settings.

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A. Regulatory Requirements for Effectiveness

53 54 The requirement that new drugs show effectiveness is based on a 1962 amendment to the Federal 55 Food, Drug, and Cosmetic Act. This law requires substantial evidence of effectiveness and 56 specifies that this evidence must be derived from adequate and well-controlled clinical 57 investigations. Clinical benefits that have supported drug approval have included important 58 clinical outcomes (e.g., increased survival, symptomatic improvement) but have also included 59 effects on established surrogate endpoints (e.g., blood pressure or serum cholesterol).

60

61 In 1992, the accelerated approval regulations (21 CFR part 314, subpart H and 21 CFR part 601, subpart E) allowed use of additional endpoints for approval of drugs or biological products that 62 63 are intended to treat serious or life-threatening diseases and that either demonstrate an 64 improvement over available therapy or provide therapy where none exists. In this setting, the 65 FDA may grant approval based on an effect on a surrogate endpoint that is *reasonably likely* to predict clinical benefit ("based on epidemiologic, therapeutic, pathophysiologic, or other 66 67 evidence"). These surrogates are less well-established than surrogates in regular use, such as 68 blood pressure or cholesterol for cardiovascular disease. A drug is approved under the 69 accelerated approval regulations on condition that the manufacturer conduct clinical studies to 70 verify and describe the actual clinical benefit. If the postmarketing studies fail to demonstrate 71 clinical benefit or if the applicant does not demonstrate due diligence in conducting the required 72 studies, the drug may be removed from the market under an expedited process. From December 73 1992 to June 2004, 22 cancer drug applications were approved under the accelerated approval 74 regulations. In the following discussion, we will use the term *regular approval* to designate the 75 longstanding route of drug approval based on demonstrating clinical benefit to distinguish it 76 from accelerated approval associated with use of a surrogate endpoint that is reasonably likely to 77 predict benefit.

78

79 The nature of evidence to support drug approval, including the preferred number of clinical

80 trials, is discussed in general FDA guidance documents. In most cases, the FDA has

- 81 recommended at least two well-controlled clinical trials. In some cases, the FDA has found that
- 82 evidence from a single trial was sufficient, but generally only in cases in which a single

multicenter study provided highly reliable and statistically strong evidence of an important
clinical benefit, such as an effect on survival, and in which confirmation of the result in a second
trial would be practically or ethically impossible.⁴ For drugs approved for treatment of patients
with a specific stage of a particular malignancy, evidence from one trial may be sufficient to
support an efficacy supplement for treatment of a different stage of the same cancer.⁵

88 89

B. Endpoints Supporting Past Approvals in Oncology

90 91 For regular approval, it is critical that the sponsor show direct evidence of clinical benefit or 92 improvement in an established surrogate for clinical benefit. In oncology, survival is the gold 93 standard for clinical benefit, but the FDA has accepted other endpoints for cancer drug approval. 94 Indeed, in the 1970s the FDA usually approved cancer drugs based on objective response rate 95 (ORR), determined by tumor assessments from radiologic tests or physical exam. In the early 96 1980s, after discussion with the ODAC, the FDA determined that it would be more appropriate 97 for cancer drug approval to be based on more direct evidence of clinical benefit, such as 98 improvement in survival or in a patient's quality of life (QOL), improved physical functioning,

99 or improved tumor-related symptoms — benefits not always predicted by ORR.

100

101 Over the next decade, several endpoints were used as surrogates for benefit. Improvement in

102 disease-free survival supported drug approval in selected surgical adjuvant settings (when a large

103 proportion of patients had cancer symptoms at the time of recurrence). Durable complete

104 response was considered an acceptable endpoint in testicular cancer and acute leukemia (a de

105 facto improvement in survival because the untreated conditions were quickly lethal) and in some 106 chronic leukemias and lymphomas (where it was clear that remission would lead to less

107 infection, bleeding, and blood product support). The FDA has also considered that a very high

108 ORR alone might sometimes support regular approval, but that response duration, relief of

109 tumor-related symptoms, and drug toxicity should also be considered (O'Shaughnessy and

110 Wittes et al., 1991, Commentary Concerning Demonstration of Safety and Efficacy of

111 Investigational Anticancer Agents in Clinical Trials, J Clin Oncol 9:2225-2232). ORR has been

an especially important endpoint for the less toxic drugs, such as the hormonal drugs for breast

113 cancer, where improvement in this endpoint has been the basis for regular approval.

114 Improvement in tumor-related symptoms in conjunction with an improved ORR and an adequate

response duration supported approval in several clinical settings.

116

117 In the last decade, in addition to its limited role in regular approval, ORR has been the primary

118 surrogate endpoint used to support cancer drug accelerated approval for several reasons. First,

119 ORR is directly attributable to drug effect (tumors rarely shrink spontaneously and, therefore,

120 ORR can be accurately assessed in single-arm studies). Second, tumor response is widely

121 accepted as relevant by oncologists and has a long-accepted role in guiding cancer treatment.

Finally, if the ORR is high enough and the responses are of sufficient duration, ORR does indeed

seem *reasonably likely* to predict clinical benefit.

⁴ See guidance for industry *Providing Clinical Evidence of Effectiveness for Human Drug and Biological Products* (http://www.fda.gov/cder/guidance/index.htm)

⁵ See guidance for industry *FDA Approval of New Cancer Treatment Uses for Marketed Drug and Biological Products* (http://www.fda.gov/cder/guidance/index.htm)

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Drugs approved under accelerated approval regulations must provide a benefit over available
 therapy. To satisfy this requirement, many sponsors have designed single-arm studies in patients

- 127 with refractory tumors where, by definition, no available therapy exists.
- 128 129

130 III. GENERAL ENDPOINT CONSIDERATIONS131

132 The following is an overview of general issues in cancer drug development. A discussion of 133 commonly used cancer endpoints is followed by a discussion of pertinent issues in cancer 134 clinical trial design using these endpoints. Future guidance documents will discuss these issues 135 in more detail with regard to specific treatment indications. Endpoints that will be discussed 136 include overall survival, endpoints based on tumor assessments (e.g., disease-free survival, ORR, 137 time to progression, progression-free survival, time to treatment failure), and endpoints based on 138 symptom assessment. A comparison of important endpoints in cancer drug approval is provided 139 in Table 1. Many of the issues relating to the proper analysis of efficacy endpoints are addressed in general FDA guidance documents.⁶ Issues that commonly arise in oncology applications are 140 discussed in this guidance. 141

142

Endpoint	Regulatory Nature of Evidence	Assessment	Some Advantages	Some Disadvantages
Overall Survival	Clinical benefit	 Randomized studies needed Blinding not essential 	 Universally accepted direct measure of benefit Easily measured Precisely measured 	 Requires larger studies Requires longer studies Potentially affected by crossover therapy Does not capture symptom benefit Includes noncancer deaths
Disease- Free Survival	Surrogate for accelerated approval or regular approval*	 Randomized studies needed Blinding preferred 	 Considered to be clinical benefit by some Needs fewer patients and shorter studies than survival 	 Not a validated survival surrogate in most settings Not precisely measured; subject to assessment bias Various definitions exist

143 **Table 1. A Comparison of Important Cancer Approval Endpoints**

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*Adequacy as a surrogate endpoint for accelerated approval or regular approval is highly dependent upon other factors such as effect size, effect duration, and benefits of other available therapy. See text for details.

continued

⁶ See ICH guidance for industry *E9 Statistical Principles for Clinical Trials* (http://www.fda.gov/cder/guidance/index.htm)

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Endpoint	Regulatory Nature of Evidence	Assessment	Some Advantages	Some Disadvantages
Objective Response Rate (ORR)	Surrogate for accelerated approval or regular approval*	 Single-arm or randomized studies can be used Blinding preferred in comparative studies 	Can be assessed in single-arm studies	 Not a direct measure of benefit Usually reflects drug activity in a minority of patients Data are moderately complex compared to survival
Complete Response (CR)	Surrogate for accelerated approval or regular approval*	 Single-arm or randomized studies can be used Blinding preferred in comparative studies 	 Durable CRs represent obvious benefit in some settings (see text) Can be assessed in single-arm studies 	 Few drugs produce high rates of CR Data are moderately complex compared to survival
Progression Free Survival (PFS)	Surrogate for accelerated approval or regular approval*	 Randomized studies needed Blinding preferred Blinded review recommended 	 Activity measured in responding and stable tumors Usually assessed prior to change in therapy Less missing data than for symptom endpoints Assessed earlier and in smaller studies compared with survival 	 Various definitions exist Not a direct measure of benefit Not a validated survival surrogate Not precisely measured compared with survival Is subject to assessment bias Frequent radiologic studies are needed Data are voluminous and complex compared to survival
Symptom Endpoints	Clinical benefit	 Usually needs randomized blinded studies (unless endpoints have an objective component and effects are large — see text) 	• Direct measure of benefit	 Blinding is often difficult in oncology trials Missing data are common Few instruments are validated for measuring cancer-specific symptoms Data are voluminous and complex compared to survival

148	Table 1,	continued
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*Adequacy as a surrogate endpoint for accelerated approval or regular approval is highly dependent upon other factors such as effect size, effect duration, and benefits of other available therapy. See text for details.

150 151 Abbreviations: complete response (CR); objective response rate (ORR); progression-free survival (PFS).

152

Overall Survival A.

153 154

155 Overall survival is defined as the time from randomization until death from any cause, and is

156 measured in the intent to treat (ITT) population. Survival is the most reliable cancer endpoint,

157 and when studies can be conducted to adequately assess it, it is usually the preferred endpoint.

An improvement in survival is of unquestioned clinical benefit. The endpoint is precise and easy 158

159 to measure, documented by the date of death. Bias is not a factor in endpoint measurement.

160

- 161 Overall survival almost always needs to be evaluated in randomized controlled studies.
- 162 Historically controlled data are seldom reliable for time-dependent endpoints such as overall
- 163 survival unless treatment effects are extreme (e.g., acute leukemia, testicular cancer). Apparent
- 164 differences in outcome between historical controls and current treatment groups can arise from
- 165 differences other than drug treatment, including patient selection, improved imaging techniques
- 166 (which can alter tumor staging and prognosis), or improved supportive care. Randomized
- studies minimize the effect of such differences by allowing a comparison of outcomes in patient groups where such factors should be similar. Demonstration of a statistically significant
- 169 improvement in overall survival is usually considered to be clinically significant, and has often
- 170 supported new drug approval.
- 171

172 Criticisms of survival as an endpoint stem not from doubts about the worth of a proven survival

benefit, but from difficulties in performing studies large enough or long enough to detect a

- survival improvement, difficulties in determining a drug's effect on survival because of the
- 175 confounding effects of subsequent cancer therapy, or a concern that the drug may be effective in
- 176 only a small fraction of those treated, making it difficult to see an effect on survival in the whole 177 population.
- 178
- 179

B. Endpoints Based on Tumor Assessments

- In this section we discuss several endpoints that are based on tumor assessments and are
 therefore unique to oncology. These endpoints include disease-free survival, objective response
 rate, time to progression, progression-free survival, and time to treatment failure. The data
- 184 collection and analysis of all time-dependent endpoints is complex, particularly when the
- assessments are indirect and based on calculations and estimates as is the case for tumor
- 186 measurements. The discussion of progression-free survival data collection and analysis is
- 187 particularly complex and is supplemented by tables in Appendix 3 of this guidance.
- 188
- 189 Selection of tumor-assessment endpoints for efficacy trials should include two judgments. First, 190 will the endpoint support accelerated approval (is the endpoint a surrogate reasonably likely to
- 191 predict clinical benefit and does the drug provide an advantage over available therapy) or regular
- approval (is it an established and/or validated surrogate for, or a direct measure of, clinical
- benefit)? Second, will the results be reliable, given the potential for uncertainty or bias in tumor
- endpoint assessments? Drug applications using studies that rely on tumor measurement based
- 195 endpoint assessments. Drug appreations using studies that rely on tamor measurement based 195 endpoints as sole evidence of efficacy should generally provide confirmatory evidence from a
- second trial. Both the precision and the clinical meaning of endpoints based on tumor
- 197 assessments can vary in different cancer settings. For instance, response rate determinations in
- 198 malignant mesothelioma and pancreatic cancer are often unreliable because of the difficulty in
- 199 measuring these tumors with currently available imaging modalities.
- 200
- 201 When the primary study endpoint for drug approval is based on tumor measurements (e.g.,
- 202 progression-free survival or ORR), it is recommended that tumor endpoint assessments generally
- 203 be verified by central reviewers blinded to study treatment (see Appendix 4), especially when the
- study itself cannot be blinded. Although the FDA will generally not ask that all tumor images be
- submitted with the marketing application, it may need to audit a sample of the scans to verify the

206 central review process. In all cases, we recommend submitting primary electronic data
 207 documenting tumor measurements and assessments.⁷ Additional details regarding data
 208 collection are listed in Appendix 1.

209 210

1. Disease-Free Survival

- 211 212 Disease-free survival (DFS) is usually defined as the time from randomization until recurrence of 213 tumor or death from any cause. Although DFS can also be an important endpoint when a large 214 percentage of patients achieve complete responses with chemotherapy, the most frequent use of 215 this endpoint is in the adjuvant setting after definitive surgery or radiotherapy. In either of these 216 settings, DFS has special meaning to patients because until a recurrence occurs, a patient can 217 hope for cure. Whereas overall survival is the standard endpoint for most adjuvant settings, DFS 218 has been the primary basis of approval for hormonal therapy after initial surgery for breast 219 cancer. An important consideration is whether prolongation of DFS represents intrinsic benefit 220 or only a potential surrogate for survival prolongation. In December 2003, the consensus of the 221 ODAC was that prolongation of DFS represented clinical benefit, but that the magnitude of this 222 benefit should be carefully weighed against the toxicity of adjuvant treatment, particularly as 223 measured by effects on patient function. In May 2004, the ODAC recommended that DFS be 224 considered an acceptable endpoint for colon cancer drugs in the surgical adjuvant setting, provided certain conditions were met.⁸ Additional cancer-specific guidances will address the 225 acceptability of DFS in other cancer settings. 226
- 227

Important considerations in evaluating DFS as a potential endpoint include the estimated size of 228 229 the treatment effect, proven benefits of standard therapies, and details of trial design. For 230 instance, when a new drug is compared to a control drug that is known to improve overall 231 survival, an important consideration is whether the DFS of the new drug is superior to, or only 232 noninferior to, the control. Clearly, proof of superiority with regard to a surrogate endpoint is 233 more persuasive than a demonstration of noninferiority. Furthermore, relying on a conclusion of 234 noninferiority based on a surrogate endpoint to support a conclusion of noninferiority with 235 respect to the definitive endpoint is problematic. Another critical issue is whether the duration of 236 study follow-up is adequate to evaluate the durability of the DFS benefit. 237

We suggest that the protocol carefully detail both the definition of DFS and the schedule for 238 239 follow-up studies and visits. Unscheduled assessments can occur for many reasons (including 240 tumor-related symptoms, drug toxicity, anxiety), and differences between study arms in the 241 frequency or reason for unscheduled assessments is likely to introduce bias. This potential bias 242 can be minimized by blinding patients and investigators to the treatment assignments if feasible. 243 The potential effects of bias due to unscheduled assessments can be evaluated by comparing their 244 frequency between treatment arms and by performing statistical analyses that assign events from 245 unscheduled visits to the time of the next scheduled visit.

⁷ See guidance for industry *Cancer Drug and Biological Products* — *Clinical Data in Marketing Applications* (http://www.fda.gov/cder/guidance/index.htm)

⁸ Transcripts are available at http://www.fda.gov/cder/drug/cancer_endpoints/default.htm.

247 Another issue in defining DFS is whether deaths occurring without prior documentation of tumor 248 progression should be scored as DFS events (disease recurrences) or should be censored in the 249 statistical analysis. All methods for statistical analysis of deaths have limitations. The approach 250 that seems less prone to introducing bias is to consider all deaths as recurrences. Limitations of 251 this approach are a potential decrease in statistical power of the study (by *diluting* the cancer-252 related events with deaths not related to cancer) and a potential to falsely prolong the DFS 253 estimates in patients who die after a long unobserved period. The latter could introduce bias if 254 the frequency of long-term follow-up visits is dissimilar on the study arms or if there is 255 nonrandom dropout due to toxicity. Some analyses count cancer-related deaths as DFS events 256 and censor noncancer deaths. This method has the potential for bias in the post hoc 257 determination of the cause of death. Furthermore, any method that censors patients, whether at 258 death or at the last visit, assumes that the censored patients have the same risk of recurrence as 259 noncensored patients. This critical assumption needs close examination in any setting where 260 deaths are to be censored. In settings where deaths due to causes other than cancer are common 261 (e.g., studies of patients with early metastatic prostate cancer), censoring deaths can be 262 appropriate.

263 264

2. Objective Response Rate

265 ORR is the proportion of patients with tumor shrinkage of a predefined amount lasting for a 266 267 predefined minimum period of time. Response duration is usually measured from the time of 268 initial response until documented tumor progression. The FDA has generally defined ORR as 269 the sum of partial responses plus complete responses. When defined in this manner, ORR is a 270 measure of drug antitumor activity even in a single-arm study. Some sponsors have proposed 271 including stable disease as a component of ORR; however, evaluating drug effects based on the 272 stable disease rate generally involves comparison to a randomized concurrent control. Also, 273 stable disease incorporates components of time to progression or progression-free survival, 274 which can be captured in a separate measurement. A variety of response criteria have been 275 considered appropriate, including the RECIST criteria (Therasse and Arbuck et al., 2000, New 276 Guidelines to Evaluate Response to Treatment in Solid Tumors, J Natl Cancer Inst, 92:205-16). 277 Important issues for determining the clinical and regulatory significance of ORR include 278 response duration, the percentage of complete responses, the toxicity of treatment, and associated 279 improvement in tumor-related symptoms. These issues, in addition to an assessment of benefits 280 of existing therapies, determine whether ORR will support marketing authorization, either for 281 regular approval (as a full surrogate for clinical benefit) or for accelerated approval (as a 282 reasonably likely surrogate).

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284 It is important that criteria for response and progression be detailed in the protocol, and data 285 should be carefully and completely collected at intervals specified in the protocol.

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3. Time to Progression and Progression-Free Survival

In the past, time to progression (TTP) (the time from randomization until objective tumor

290 progression) and progression-free survival (PFS) (the time from randomization until objective 291 tumor progression or death) have seldom served as primary endpoints for drug approval. Time

to symptomatic progression, which would represent a clear clinical benefit, is infrequently

293 assessed but would be a credible endpoint of a well-conducted (generally blinded) trial. In 294 December 2003, the ODAC discussed both potential roles of TTP and PFS in cancer drug approval and the committee's preference for PFS versus TTP.⁹ The ODAC suggested relying on 295 these endpoints in selected clinical situations, such as diseases with low complete response rates 296 297 or when documentation of a survival benefit in clinical trials can be difficult. In settings where 298 most patients are symptomatic, the ODAC preferred measuring tumor response and symptom 299 benefit. The definition of tumor progression varies widely; therefore, it is important that it be 300 carefully detailed in the protocol.

301 302

303

a. TTP vs. PFS

304 The ODAC consensus was that PFS is a better predictor of clinical benefit than TTP and thus 305 preferable as a drug approval endpoint when used as a surrogate for clinical benefit (rather than 306 just as an indicator of antitumor activity) because PFS includes deaths. Unanticipated effects of 307 drugs on survival would thus be included in the endpoint. In the analysis of TTP, deaths are 308 censored, either at the time of death or at an earlier visit. This approach is questionable because 309 it can represent *informative censoring* (i.e., there may be a nonrandom pattern of loss from the 310 study). It seems unlikely in most cancer settings that patient deaths are randomly related to 311 tumor progression (e.g., it is likely that some deaths result from complications of undocumented 312 cancer progression). Therefore, in most settings PFS is the preferred regulatory endpoint. In 313 settings where most deaths are due to causes other than cancer, however, TTP can be an 314 appropriate endpoint.

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- 316 317

b. PFS as an endpoint to support drug approval

318 Some advantages and disadvantages of using PFS as an endpoint to support cancer drug approval 319 are listed in Table 1. Conceptually, PFS has desirable qualities of a surrogate endpoint because it 320 reflects tumor growth (a phenomenon likely to be on the causal pathway for cancer-associated 321 morbidity and death), can be assessed prior to demonstration of a survival benefit, and is not 322 subject to the potential confounding impact of subsequent therapy (unless worsening of a blood 323 marker leads to a change in treatment prior to progression). Moreover, an effect on PFS occurs 324 earlier than an effect on survival, so that a given advantage, say a median improvement of 3 325 months, represents a larger (and thus more detectable) hazard ratio improvement than would a 3-326 month median survival benefit occurring later. The formal validation of PFS as a surrogate for 327 survival for the many different malignancies that exist, however, would be difficult. Data are 328 usually insufficient to allow a robust evaluation of the correlation between effects on survival 329 and PFS. Oncology trials are often small, and proven survival benefits of existing drugs are 330 generally modest. The role of PFS as an endpoint to support licensing approval varies in 331 different cancer settings. In some settings PFS prolongation might be an accepted surrogate 332 endpoint for clinical benefit to support regular approval, and in others it may be a surrogate 333 reasonably likely to predict benefit for accelerated approval. Important considerations will be 334 the magnitude of the effect, the toxicity profile of the treatment, and the clinical benefits and 335 toxicities of available therapies. These issues will be discussed in future guidance documents for 336 specific cancer settings. 337

⁹ Transcripts are available at http://www.fda.gov/cder/drug/cancer_endpoints/default.htm.

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338 PFS trial design issues C. 339 340 It is important that methodology for assessing, measuring, and analyzing PFS be detailed in the 341 protocol and statistical analysis plan. It is also important to carefully define tumor progression 342 criteria in the protocol. There are no standard regulatory criteria for defining progression. 343 Sponsors have used a variety of different criteria, including the RECIST criteria. The broad 344 outline presented in most published PFS criteria should be supplemented with additional details 345 in the protocol and statistical analysis plan. It is important that visits and radiological 346 assessments be symmetric on the two study arms to prevent systematic bias. When possible, 347 studies should be blinded. Blinding is particularly important when patient or investigator 348 assessments are included as components of the progression endpoint. It is important that the 349 FDA and the sponsor agree prospectively on the protocol, data to be recorded on the case report 350 form, statistical analysis plan (including analysis of missing data and censoring methods), and, if 351 applicable, the operating procedures of an independent endpoint review committee (discussed in 352 Appendix 4). The effect of follow-up visit frequency has been debated. Frequent regular 353 assessments, depending on the type and stage of cancer, ensure that most progression events will 354 be detected on radiologic scans rather than as symptomatic events. This approach increases the 355 expense and difficulty of the study, including an increased data collection burden on the 356 investigator and an increased number of scans for patients, and may not mirror clinical practice 357 standards. 358

359 360

d. Analysis of PFS

361 The analysis of PFS is complicated by missing data. It is important that the protocol specify 362 what constitutes an adequate assessment visit for each patient (i.e., a visit when all scheduled 363 tumor assessments have been done). The analysis plan should outline a comparison of the 364 adequacy of follow-up in each treatment arm and specify how incomplete or missing follow-up 365 visits will be handled with regard to censoring. For instance, if one or more assessment visits are 366 missed just prior to the progression event, to what date should the progression event be assigned? 367 It is important that the analysis plan specify the primary analysis and one or more sensitivity 368 analyses. For instance, in the previous example, the primary analysis might assign the actual 369 date of observed progression as the progression date. The sensitivity analysis might censor the 370 data at the last adequate assessment visit. Although both analyses are problematic (the best 371 solution to missing data is to have none), the conclusion is probably valid if it is supported by the 372 results of both the primary and the sensitivity analyses. Other methods could be considered if 373 adequately supported by the sponsor. The analysis plan should evaluate the number of deaths in 374 patients who have been lost to follow-up for more than a substantial (prespecified) time. An 375 imbalance in such deaths could bias the measurement of PFS, artificially prolonging PFS on the 376 arm with less adequate follow-up.

377

Because progression data can be collected from a variety of sources (including physical exams at

unscheduled visits and radiologic scans of various types) and at a variety of times, it is important

that data collection efforts for each assessment visit be limited to a specified short time interval

381 prior to the visit. When data are collected over a longer time, the question then arises: What

date should serve as the progression date or the censoring date? A common method is to assign

383 progression to the earliest observed time when an observation shows progression and to censor at

- 384 the date when the last radiologic assessment determined a lack of progression. Because this 385 method could introduce an assessment bias, especially in unblinded trials, we recommend 386 assigning the progression and censoring times to the time of the scheduled assessment visits. A 387 study of time to symptomatic progression, if conducted blindly and with few scheduled 388 assessments, in contrast, could use the actual time of observed symptom progression. The PFS 389 date based on a death, however, would be the date of death rather than the assigned visit date 390 since death ascertainment is not related to visit time and not subject to interpretation. 391 392 Appendix 3 provides a set of tables for potential analyses of PFS that could be used for primary 393 or sensitivity analyses. We recommend that plans for PFS data collection and analysis be 394 discussed with the FDA at end-of-phase 2 meetings and verified in special protocol assessments. 395 396 Future methods for assessing progression e. 397 398 In the future, it is important that other methods of progression assessment be evaluated as 399 potential surrogate endpoints for regular approval or accelerated approval. One proposed 400 method (not used to date) is the single time point assessment which could decrease the 401 complexity of progression assessment and eliminate time-dependent assessment bias. In the 402 single time point analysis, progression would be assessed at baseline and at one prespecified time 403 after randomization. If patients progress prior to the specified time, radiologic scans could 404 document progression and the patient could go off-study. All other patients would have a 405 detailed radiologic evaluation at the prespecified follow-up time. The statistical analysis could 406 compare the proportions of patients on each study arm with progression on or before the 407 prespecified time after randomization. Potential problems with this approach are decreased 408 statistical power, potential for missing a small benefit at a time different from the prespecified 409 time, and lack of information regarding the relationship between the single time point analysis 410 and the familiar endpoints of progression-free survival and overall survival. Although this 411 approach could provide some advantages and decrease assessment bias, study dropouts prior to 412 progression could present the same difficulty as they do for all progression endpoints. Settings 413 in which further evaluation of this approach seems warranted are those where a significant and 414 durable effect on progression-free survival is expected and where complete progression-free 415 survival data collection seems impossible or impractical.
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4. Time to Treatment Failure

419 Time to treatment failure (TTF) is a composite endpoint measuring time from randomization to 420 discontinuation of treatment for any reason (including progression of disease, treatment toxicity, 421 and death). Defined that way, TTF is not recommended as an endpoint for drug approval 422 because it combines efficacy and toxicity measures. For example, suppose the standard 423 comparator (Drug A) provides a known survival benefit, but only at the cost of considerable 424 toxicity with many patients leaving therapy because of that toxicity. A nontoxic investigational 425 drug (Drug B) could have a significantly longer TTF than Drug A solely because it caused fewer 426 toxic dropouts. These data alone could not support drug approval because they would not 427 demonstrate that Drug B is effective. Drug approval would require a demonstration of Drug B 428 efficacy, such as a survival improvement or other clinical benefit. 429

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C. Endpoints Involving Symptom Assessment

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432 Symptomatic improvement has always been considered a clinical benefit, and many FDA cancer 433 drug approvals have used patient symptom assessments and/or physical signs thought to 434 represent symptomatic improvement (e.g., weight gain, decreased effusion) as the primary evidence of effectiveness. To date, broader measures of health-related quality of life (HRQL 435 436 instruments) have not served this role. HRQL is discussed in a separate FDA draft guidance on patient-reported outcomes (PRO).¹⁰ The FDA has relied on symptom scores, signs, and 437 symptoms representing obvious benefit (e.g., decreased esophageal obstruction, fewer bone 438 439 fractures, reduced size and number of skin lesions, physician actions [need for radiation therapy 440 in response to painful bone metastases], physician assessments of performance status, and 441 patient-reported assessments of symptom scales). Relying on such evidence of clinical benefit as 442 the basis for approval has allowed the FDA to approve cancer drugs earlier than if demonstration 443 of a survival benefit had been required. It seems self-evident that cancer patients will be in most 444 cases the best source for determining effects on patient symptoms, so that PRO instruments seem 445 most appropriate. Formal PRO instruments can be designed that focus on specific symptoms 446 (e.g., a pain scale) or on a broader array of physical, emotional, and activity measures. 447

448 The use of improvement of signs and symptoms or QOL assessments as primary endpoints to

support cancer drug approval requires discrimination between tumor symptoms and drug
 toxicity, especially when evidence is based on comparison to a toxic active control. This poses

451 particular problems for general HRQL scales, which, by definition, are multidimensional scales

452 including elements other than physical problems. An apparent effectiveness advantage of one

453 drug over another measured on a global HRQL instrument might simply indicate less toxicity of

454 one product or regimen versus the other, a matter of interest but not an effectiveness measure.

455 Morbidity endpoints used to date for cancer drug approvals have possessed *face validity* (value 456 obvious to patients and physicians, for example, an endpoint based on functional measures such

457 as the ability to swallow solids, liquids, or nothing) and have not measured benefit and toxicity

- 458 on the same scale.
- 459 460

1. Specific Symptom Endpoints

461 462 One endpoint the FDA has suggested to sponsors is *time to progression of cancer symptoms*, an 463 endpoint similar to time to progression. This endpoint would be a direct measure of clinical 464 benefit rather than a potential surrogate. Sponsors have cited several problems with this approach. First, because few cancer trials are blinded, assessments can be biased and therefore 465 466 unreliable. Another problem is the usual delay between tumor progression and the onset of 467 cancer symptoms. Often alternative treatments are begun before reaching the symptom endpoint. 468 which can confound the results. Many cancer trials are performed in patients with little prior 469 exposure to chemotherapy and who usually have minimal cancer symptoms. Finally, it can

sometimes be difficult to differentiate tumor symptoms from drug toxicity, a problem noted in

¹⁰ The draft guidance for industry *Patient-Reported Outcome Measures: Use in Medical Product Development to Support Claims* is currently being developed and is expected to publish in the summer of 2005. When final, this guidance will represent the FDA's current thinking on this topic. For the most recent version of a CDER or CBER guidance, check the CDER guidance Web page at http://www.fda.gov/cder/guidance/index.htm and the CBER Web page at http://www.fda.gov/cber/guidance/index.htm.

- 471 discussions of time to treatment failure and HRQL. Time to progression of symptoms and time to 472 onset of symptoms can be reasonable endpoints in cancer settings where treatment can be 473 blinded, most progressing patients are symptomatic, no effective therapy exists, and less frequent 474 radiologic follow-up is appropriate. Symptom data should be carefully collected using a 475 validated instrument according to a schedule detailed in the protocol. 476 477 A composite symptom endpoint can be appropriate when the benefit of a drug is multifaceted. It 478 is important that the components of the endpoint be related and generally of similar clinical
- 479 importance. Drugs have been approved for treatment of patients with cancer metastases to the 480 skeleton based on a composite benefit endpoint consisting of one or more skeletal-related event 481 (SRE) that would be anticipated to be associated with pain and other distress. SREs are defined 482 as pathologic fractures, radiation therapy to bone, surgery to bone, and spinal cord compression. 483 Clinical Benefit Response, a composite endpoint of pain and analgesic consumption reported by 484 the patient, and performance status assessed by a physician, in part supported approval of a drug 485 to treat pancreatic cancer.
- 486
- 487 Selection of the appropriate population for study can be critical for documenting symptom 488 benefit. Patients symptomatic at study baseline can be evaluated with a categorical symptom 489 response analysis. This approach can be appropriate for diseases such as lung cancer, when most 490 patients have symptoms at diagnosis. Studies of asymptomatic patients could use a time-to-first-491 symptom analysis. Even if the patient discontinues the study drug or begins a new drug, 492 symptomatic progression could still be assessed if follow-up is continued until documentation of 493 the first symptom. This approach is worth considering but has been infrequently attempted.
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2. Problems Encountered with Symptom Data

- 497 Many problems have been encountered in the analysis of symptom data submitted to the FDA. 498 The most important problem in oncology is that few trials are blinded so that the possibility of 499 observer bias is difficult to exclude. Missing data are common and often cast doubt on study 500 conclusions. It is critically important to have frequent assessments to minimize long unobserved 501 gaps. In addition, symptom severity should be addressed, rather than providing only a binary 502 present or absent. Withdrawing treatment because of drug toxicity or tumor progression is one 503 cause of missing symptom data. Ideally, when patients stop treatment, data collection forms 504 should continue to gather information to inform the analysis. Symptom data could lead to a large 505 number of different endpoints, and prospectively defined statistical plans need to correct for 506 multiplicity if each symptom is treated as a separate endpoint.
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D. **Biomarkers**

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510 To date, evidence from biomarkers assaved from blood or body fluids has not served as primary 511 endpoints for cancer drug approval, although paraprotein levels measured in blood and urine 512 have contributed to response endpoints for myeloma. Further research is needed to establish the 513 validity of the available tests and determine whether improvements in such biomarkers are 514 reasonably likely to predict clinical benefit (accelerated approval) or are established surrogates 515 for clinical benefit (regular approval).

517 Although tumor markers are not yet used alone as a basis for marketing approval, the FDA has 518 sometimes accepted their inclusion as elements in composite endpoints. For instance, women 519 with ovarian cancer often show clinical deterioration from progression of unmeasured tumor. In 520 blinded randomized controlled trials in advanced refractory ovarian cancer, the FDA has 521 accepted use of a composite endpoint that included CA-125. The occurrence of certain clinical 522 events (a significant decrease in performance status, or bowel obstruction) coupled with marked 523 increases in CA-125 was considered progression in these patients. The use of prostate specific 524 antigen (PSA) was discussed at a recent workshop on prostate cancer endpoints. Different 525 methods of evaluating PSA as an endpoint were discussed, including PSA response, PSA slope, 526 and PSA velocity. Although the FDA has not yet accepted a PSA endpoint to support drug 527 approval, evaluation of additional data and further discussions of PSA endpoints are planned in future workshops and ODAC meetings.¹¹ 528

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IV. ENDPOINTS AND CLINICAL TRIAL DESIGN; SELECTED ISSUES

By law, the FDA must base new drug approval decisions on substantial evidence of efficacy
from "adequate and well-controlled investigations." Regulations describe the meaning of
"adequate and well-controlled investigations." Studies must allow a valid comparison to a
control and must provide a quantitative assessment of the drug's effect. (See 21 CFR 314.126.)
Below we discuss several issues related to the design of cancer trials intended to support drug
approval.

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A. Single-Arm Studies

542 The most reliable method for demonstrating efficacy is to show a statistically significant 543 improvement in a clinically meaningful endpoint in blinded randomized controlled trials. Other 544 approaches have also been successful in certain settings. In settings where there is no effective 545 therapy and where major tumor regressions can be presumed to occur infrequently in the absence 546 of treatment (a historical control), the FDA has sometimes accepted ORR and response duration 547 observed in single-arm studies as substantial evidence supporting accelerated approval or even 548 regular approval (e.g., when many complete responses were observed or when toxicity was 549 minimal or modest). In contrast to the success of this approach, evidence from historically 550 controlled trials attempting to show improvement in time-to-event endpoints such as survival. 551 time to progression, or progression-free survival have seldom been persuasive support for drug 552 approval, except when treatment provides survival outcomes that contrast markedly with 553 historical experience (e.g., testicular cancer, acute leukemias). In most cases, however, these 554 outcomes vary among study populations in ways that cannot always be predicted; for example, 555 changes in concomitant supportive care or frequency and method of tumor assessment can differ 556 by location or change over time. Consequently, comparisons involving these time-to-event 557 endpoints generally need a concurrent control (preferably in a randomized trial), unless, as noted, 558 the effect is very large.

¹¹ Transcripts are available at http://www.fda.gov/cder/drug/cancer_endpoints/default.htm.

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561 B. Studies Designed to Demonstrate Noninferiority

- 562 563 The goal of noninferiority (NI) trials is to demonstrate the effectiveness of a new drug showing 564 that it is not less effective, by a predefined amount, than a standard regimen known to have the 565 effect being investigated (Temple and Ellenberg, 2000, Placebo-Controlled Trials and Active-Control Trials in the Evaluation of New Treatments, Part 1: Ethical and Scientific Issues, Ann 566 Intern Med, 2000 Sep 19; 133(6):455-63).¹² The difference to be ruled out, the *noninferiority* 567 568 margin, cannot be larger than the effect of the control drug in the new study. As that effect is not 569 measured (the new study does not have a no-treatment arm), the effect must be assumed based 570 on the previous studies of the control drug that documented its effect. If the new drug is inferior 571 by more than the noninferiority margin, it would have no effect at all. In most cases the NI 572 margin is not set at the control drug's full effect, but at some fraction of it (e.g., 50 percent), so 573 that the study seeks to show that at least 50 percent of the control drug effect is preserved.
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575 There are multiple difficulties with NI trials. NI trials rely on historical data to establish the 576 expected size of treatment effect of the active control. In many situations adequate historical 577 data for the control do not exist. Moreover, a critical assumption is that the treatment effect of 578 the active control that was observed historically will also be observed in the current population in 579 the new study. This assumption is difficult to support, as results of trials are almost never 580 identical (although one can evaluate control regimen response rates in the historical and NI trial 581 populations as some measure of comparability). Optimally, the estimated size of the treatment 582 effect of the active control would be based on a comprehensive meta-analysis of historical 583 studies that reproducibly demonstrate the effectiveness, compared to no treatment, of the control 584 agent. In the oncology setting, however, information is often lacking on effects compared to a 585 no-treatment control. The variability in the meta-analysis will be reflected in the choice of the 586 noninferiority margin. But there may be little data from randomized controlled trials available to 587 estimate the treatment effect and thus no basis for estimating the control treatment effect. 588 Furthermore, subsequent events in the trial, especially crossover from the control, can invalidate 589 NI survival analyses (producing a bias toward a showing of no difference). NI designs generally 590 require many patients in order to provide meaningful results. Given the complex issues 591 involved, we strongly recommend that sponsors designing noninferiority trials consult early with 592 the FDA. Because of the difficulties with the design, conduct, and analysis of NI trials, a single 593 NI trial seldom provides sufficient evidence of efficacy to support drug approval. 594

595 When the new treatment has a different toxicity profile from available treatments, it may be 596 possible to *design around* the NI study problem by conducting an *add-on* study, adding new drug 597 or placebo/no treatment to the standard therapy. This will not be possible if the goal is to show a 598 new treatment to be less toxic than existing therapy (but still effective). In this case the NI 599 design is unavoidable in order to demonstrate that the survival benefit of the standard drug is 600 retained by the experimental drug. If the standard drug is associated with only a small proven 601 survival benefit, however, interpretation of an NI study is difficult or impossible. Moreover, the 602 size of such NI trials can be prohibitively large.

¹² See ICH guidance for industry *E10 Choice of Control Group and Related Issues in Clinical Trials* (http://www.fda.gov/cder/guidance/index.htm)

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604 С. No Treatment or Placebo Control

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606 Giving no anticancer drug treatment to patients in the control arm of a cancer study is often 607 considered unethical, but, in some settings, it can be acceptable. For instance, in early stage 608 cancer when standard practice is to give no treatment, comparison of a new agent to a no-609 treatment control would be acceptable. This approach would not be an ethical problem in the so-610 called *add-on* design, when all patients receive standard treatment plus either no additional 611 treatment or the experimental drug. Using a control group that receives only best supportive care 612 is acceptable in an advanced refractory setting where there is no effective therapy. Placebos 613 (identical appearing inactive controls) are generally preferred to no-treatment controls because 614 they permit blinding. With many cytotoxic cancer drugs, blinding may not be feasible because 615 of a relatively high rate of recognizable toxicities, but newer interventions, many of them much 616 less toxic, are increasingly being studied in blinded trials.

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D. **Isolating Drug Effect in Combinations**

619 620 Because marketing approval is usually for a single drug product rather than for a drug 621 combination, clinical trials supporting regulatory approval need to isolate the effectiveness of the 622 proposed agent. Evidence is needed showing not only the effectiveness of the regimen but also 623 establishing the contribution of the new drug to that regimen. One way to demonstrate the 624 individual contribution of a new drug in a regimen is using the *add-on* design previously 625 discussed. Sometimes the clinical effects seen in early phases of development can be used to 626 establish the contribution of a drug to a drug regimen, particularly if the combination is more 627 effective than any of the individual components. We recommend discussing these issues with 628 the FDA at end-of-phase 1 or end-of-phase 2 meetings.

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E. **Trial Designs for Radiotherapy Protectants and Chemotherapy Protectants**

632 Radiotherapy protectants and chemotherapy protectants are drugs designed to ameliorate the 633 toxicities of radiotherapy or chemotherapy. Trials to evaluate these agents usually have two 634 objectives. The first is to assess whether the protecting drug achieves its intended purpose of 635 ameliorating the cancer treatment toxicity. Unless the mechanism of protection is clearly 636 unrelated to the mechanism of antitumor activity (e.g., antiemetic agents which ameliorate 637 nausea via central nervous system receptors), a second trial objective is to determine whether 638 anticancer efficacy is compromised by the protectant. Because the comparison of antitumor 639 activity between the two arms of the trial is a noninferiority comparison, a large number of 640 patients may be required to achieve this objective. Generally, a second study is needed to 641 confirm the findings. A critical question for the future is whether, in such cases where the same 642 drug is studied in both arms, ORR should be considered a sufficient endpoint for comparing drug 643 activity and benefit.

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646 V. SUMMARY AND CONCLUSION 647

648 Although general principles outlined in this guidance should help sponsors select endpoints for 649 marketing applications, we recommend that sponsors meet with the FDA before submitting

- 650 protocols intended to support NDA or BLA marketing applications. The FDA will ensure that
- these meetings include a multidisciplinary FDA team of oncologists, statisticians, clinical
- barmacologists, and often external expert consultants. Sponsors may submit protocols after
- 653 these meetings and request a special protocol assessment that provides the acceptability of
- endpoints and protocol design to support drug marketing applications.¹³ Ultimately, of course,
- 655 marketing approval will depend not only on the design of a single trial, but on FDA review of the
- results and data from all studies in the drug marketing application.

¹³ See guidance for industry Special Protocol Assessment (http://www.fda.gov/cder/guidance/index.htm)

658 659	APPENDIX 1: THE COLLECTION OF TUMOR MEASUREMENT DATA ¹⁴
660 661 662 663	The following are important considerations for tumor measurement data. The Agency recommends that:
664 665 666	• The case report form (CRF) and electronic data document the target lesions identified during the baseline visit prior to treatment. Retrospective identification of such lesions would rarely be considered reliable.
667 668 669	• Tumor lesions are assigned a unique identifying letter or number. This allows differentiating among multiple tumors occurring at one anatomic site and matching of tumors measured at baseline and tumors measured during follow-up.
670 671 672	• A mechanism ensures complete collection of data at critical times during follow-up. It is important that the CRF ensures that all target lesions are assessed at each follow-up visit and that all required follow-up tests are done with the same imaging/measuring method.
673 674	 The CRF contains data fields that indicate whether scans were performed at each visit. A zero is recorded when a lesion has completely resolved. Otherwise, disappearance of a lease a differentiated from a missing value.
675 676 677	 Follow-up tests allow timely detection of new lesions both at initial and new sites of disease. It is important that the occurrence of and location of new lesions be recorded in the CRF and
678 679 680	the submitted electronic data.

¹⁴ *Tumor data* in this section refers to data in SAS transport files, not images. Images are not generally submitted to the NDA/BLA, but may be audited by the FDA during the review process.

681 **APPENDIX 2:** 682 **ISSUES TO CONSIDER IN PFS ANALYSIS** 683 684 The protocol and statistical analysis plan (SAP) of a study should detail the primary analysis of progression-free survival (PFS). This includes a detailed description of the endpoint, acceptable 685 686 modalities for evaluating tumors, and procedures for minimizing bias when determining 687 progression status, such as procedures for an independent endpoints review committee. It is 688 important that one or two secondary analyses be specified to evaluate anticipated problems in 689 trial conduct and to assess whether results are robust. The following are several important 690 factors to consider. 691 692 **Definition of progression date.** Survival analyses use the exact date of death. In analyses 693 of PFS, however, the exact progression date is unknown. The following are two methods for 694 defining the recorded progression date (PDate) used for PFS analysis. 695 696 1. One approach assigns PDate to the first time at which progression can be declared: 697 For progression based on a new lesion, the PDate is the date of the first observation 698 that detects the new lesion. 699 . For progression based on the sum of target lesion measurements, PDate is the date of 700 the last observation or radiologic assessment of target lesions (if multiple assessments 701 are done at different times). 702 This approach can introduce between-arm bias if radiologic assessments are done earlier 703 or more frequently in one treatment arm. 704 705 2. A second approach assigns the PDate to the date of the scheduled clinic visit immediately 706 after all radiologic assessments (which collectively document progression) have been 707 done. Although this approach provides a less accurate estimate of the true date of 708 progression, the error should be symmetrically distributed between arms, and between-709 arm bias is minimized. 710 711 **Definition of censoring date.** Censoring dates are defined in patients with no documented • 712 progression prior to data cutoff or dropout. In these patients, the censoring date is often 713 defined as the last date on which progression status was adequately assessed. One acceptable 714 approach uses the date of the last assessment performed. However, multiple radiologic tests 715 can be evaluated in the determination of progression. A second acceptable approach uses the 716 date of the clinic visit corresponding to these radiologic assessments. 717 718 Definition of an adequate PFS evaluation. In patients with no evidence of progression, • 719 censoring for PFS often relies on the date of the last *adequate tumor assessment*. A careful 720 definition of what constitutes an adequate tumor assessment includes adequacy of target 721 lesion assessments and adequacy of radiologic tests both to evaluate nontarget lesions and to 722 search for new lesions. 723 724 Analysis of partially missing tumor data. Analysis plans should describe the method for •

Analysis of partially missing tumor data. Analysis plans should describe the method fo
 calculating progression status when data are partially missing from *adequate tumor*

assessment visits. For instance, are the values for missing target lesions to be *carriedforward*?

728 729 Completely missing tumor data. Assessment visits where no data are collected are • 730 sometimes followed by death or by assessment visits showing progression; in other cases the 731 subsequent assessment shows no progression. In the latter case, at first glance, it might seem 732 acceptable to continue the patient on study and continue monitoring for evidence of 733 progression. This approach, however, treats missing data differently depending upon 734 subsequent events and could represent informative censoring. Therefore, another possibility 735 is for the primary analysis to include data from subsequent PFS assessments when only a 736 single follow-up visit is missed but censor data when there are two or more missed visits. It 737 is important that the SAP detail primary and secondary PFS analyses to evaluate the potential 738 effect of missing data. Reasons for dropouts should be incorporated into procedures for 739 determining censoring and progression status. For instance, for the primary analysis, patients 740 going off-study for undocumented clinical progression, change of cancer treatment, or 741 decreasing performance status could be censored at the last adequate tumor assessment. The 742 secondary sensitivity analysis would include these dropouts as progression events.

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Progression of nonmeasurable disease. When appropriate, progression criteria should be described for each assessment modality (e.g., CT scan, bone scan). It is important that scans documenting progression based on nonmeasurable disease be verified by a blinded review committee and be available for verification by the FDA if needed.

Suspicious lesions. Sometimes new lesions are identified as suspicious. An algorithm should be provided for following up these lesions and for assignment of progression status at the time of analysis. For example, a radiological finding identified as suspicious at visit one might be verified as being a new tumor at visit three. It is important that the protocol or analytical plan clarify whether the progression time would be visit one or visit three.

756	APPENDIX 3:
757	EXAMPLE TABLES FOR PFS ANALYSIS
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759	As discussed in Section III.B., sensitivity analyses may be helpful in determining whether the
760	PFS analysis is robust. Different sensitivity analyses can be described in tables that specify how
761	to assign dates of progression events and dates for censoring of progression data. The following
762	three tables describe examples of three different sensitivity analyses:
763	
764	a. Table A represents a sensitivity analysis that only includes well-documented and
765	verifiable progression events. Other data are censored. In Table A the progression dates
766	are:
767	 Based only on radiologic assessments verified by an independent review committee
768	(IRC). <i>Clinical progression</i> is not considered a progression endpoint.
769	 Assigned to the first time when tumor progression was noted.
770	• The date of death when the patient is closely followed. Deaths occurring after two or
771	more missed visits, however, are censored at last visit.
772	
773	Table A. PFS 1 (includes documented progression only)

Situation	Date of Progression or Censoring	Outcome
No baseline tumor assessments	Randomization	Censored
Progression documented between	Earliest of:	Progressed
scheduled visits	 Date of radiologic assessment showing new lesion (if progression is based on new lesion); or Date of last radiologic assessment of measured lesions (if progression is based on increase in sum of measured 	
	lesions)	
No progression	Date of last radiologic assessment of measured lesions	Censored
Treatment discontinuation for undocumented progression	Date of last scan of measured lesions	Censored
Treatment discontinuation for toxicity or other reason	Date of last radiologic assessment of measured lesions	Censored
New anticancer treatment started	Date of last radiologic assessment of measured lesions	Censored
Death before first PD assessment	Date of death	Progressed
Death between adequate assessment visits	Date of death	Progressed
Death or progression after more than one missed visit	Date of last radiologic assessment of measured lesions	Censored

 Table A. PFS 1 (includes documented progression only)

- The sensitivity analysis in Table B corrects for potential bias in follow-up schedules for 776
- 777 tumor assessment by assigning the dates for censoring and events only at scheduled visit 778 dates.
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780	Table B.	PFS 2 (uniform	nrogression	and	assessment da	tes)
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Situation	Date of Progression or Censoring	Outcome
No baseline tumor assessments	Randomization	Censored
Progression documented	Date of next scheduled visit	Progressed
between scheduled visits		
No progression	Date of last visit with adequate assessment	Censored
Treatment discontinuation for	Date of last visit with adequate assessment	Censored
undocumented progression		
Treatment discontinuation for	Date of last visit with adequate assessment	Censored
toxicity or other reason		
New anticancer treatment started	Date of last visit with adequate assessment	Censored
Death before first PD assessment	Date of death	Progressed
Death between adequate	Date of death	Progressed
assessment visits		_
Death or progression after more	Date of last visit with adequate assessment	Censored
than one missed visit		

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- **b.** The sensitivity analysis in Table C evaluates PFS according to the investigator's assessment.
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785 Table C. PFS 3 (includes investigator claims)

Situation	Date of Progression or Censoring	Outcome
No baseline assessment	Randomization	Censored
Progression documented between	Next scheduled visit	Progressed
scheduled visits		
No progression	Date of last visit with adequate assessment	Censored
Investigator claim of clinical	Scheduled visit (or next scheduled visit if	Progressed
progression	between visits)	
Treatment discontinuation for	Date of last visit with adequate assessment	Censored
toxicity or other reason		
New anticancer treatment started	Date of last visit with adequate assessment	Censored
with no claim of progression		
Death before first PD assessment	Date of death	Progressed
Death between adequate	Date of death	Progressed
assessment visits or after patient		
misses one assessment visit		
Death after an extended lost-to-	Last visit with adequate assessment	Censored
follow-up time (two or more		
missed assessments)		

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APPENDIX 4: INDEPENDENT REVIEW OF TUMOR ENDPOINTS

790 791 Sponsors and the FDA need to be able to verify clinical trial results that support drug approval, 792 including ORR and progression-free survival. ORR determined in single-arm studies can be 793 verified by scrutiny of a limited number of images. However, when drug approval is based on 794 measurement of progression-free survival in a randomized study, careful planning is needed to 795 minimize bias and to allow the sponsor and the FDA to verify results. This is especially true 796 when investigators and patients cannot be blinded to treatment assignment because of drug 797 toxicities or manner of administration. An independent endpoints review committee (IRC) 798 provides a mechanism to minimize bias in interpretation of the radiologic findings and 799 independent adjudication of endpoints. We recommend that a clearly described written plan 800 outlining the IRC function and process, sometimes called an independent review charter, be 801 agreed upon with the FDA prior to study conduct. It is important that the plan describe how the 802 independence of the committee will be assured; how images will be collected, stored, 803 transported, and reviewed; how differences in image interpretation will be resolved; how clinical 804 data will be used in final endpoint interpretation; and how, if needed, images and IRC results will 805 be made available to the FDA for audit. The use of an IRC is discussed further in a draft 806 guidance for the development of medical imaging products.¹⁵

¹⁵ See draft guidance for industry *Developing Medical Imaging Drug and Biological Products, Part 3: Design, Analysis, and Interpretation of Clinical Studies.* When final, this guidance will represent the FDA's current thinking on this topic. For the most recent version of a CBER guidance, check the CBER guidance Web page at http://www.fda.gov/cber/guidelines.htm.