Second cancers after fractionated radiotherapy: Stochastic population dynamics effects

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Abstract

When ionizing radiation is used in cancer therapy it can induce second cancers in nearby organs. Mainly due to longer patient survival times, these second cancers have become of increasing concern. Estimating the risk of solid second cancers involves modeling: because of long latency times, available data is usually for older, obsolete treatment regimens. Moreover, modeling second cancers gives unique insights into human carcinogenesis, since the therapy involves administering well-characterized doses of a well-studied carcinogen, followed by long-term monitoring.

In addition to putative radiation initiation that produces pre-malignant cells, inactivation (i.e. cell killing), and subsequent cell repopulation by proliferation, can be important at the doses relevant to second cancer situations. A recent initiation/inactivation/proliferation (IIP) model characterized quantitatively the observed occurrence of second breast and lung cancers, using a deterministic cell population dynamics approach. To analyze if radiation-initiated pre-malignant clones become extinct before full repopulation can occur, we here give a stochastic version of this IIP model. Combining Monte-Carlo simulations with standard solutions for time-inhomogeneous birth-death equations, we show that repeated cycles of inactivation and repopulation, as occur during fractionated radiation therapy, can lead to distributions of pre-malignant cells per patient with variance \( \text{mean} \), even when pre-malignant clones are Poisson-distributed. Thus fewer patients would be affected, but with a higher probability, than a deterministic model, tracking average pre-malignant cell numbers, would predict. Our results are applied to data on breast cancers after radiotherapy for Hodgkin disease. The stochastic IIP analysis, unlike the deterministic one, indicates: (a) initiated, pre-malignant cells can have a growth advantage during repopulation, not just during the longer tumor latency period that follows; (b) weekend treatment gaps during radiotherapy, apart from decreasing the probability of eradicating the primary cancer, substantially increase the risk of later second cancers.

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1. Introduction

Ionizing radiation is a carcinogen as well as an agent for killing cells. When radiotherapy is used to eradicate a tumor, the radiation can cause second cancers, e.g. in organs adjacent to the tumor (reviewed in Curtis et al., 2006; Hall, 2006; Little, 2001; Ron, 2003). With screening resulting in patients being treated at younger ages, and with increasing patient survival times, second cancers are becoming of increasing concern (Travis et al., 2006). The long time lag between radiotherapy and second cancer incidence means that few direct data, or none, are available on second cancers induced by recently introduced radiation conditions. Thus, using the IIP model to predict second cancer incidence can be important, particularly for high doses and long latency periods.
treatment modalities, so that model-based predictions are important.

1.1. Models of radiation carcinogenesis used in applied risk estimation

Radiation carcinogenesis, in this second cancer situation and in general, is a complex process. But it is important to have a well defined, consensus, quantitative method to get some estimate of the cancer risk. In many practical applications (e.g. Bennett et al., 2004; Brenner et al., 2003; Land et al., 2003; Preston et al., 2003; Walsh et al., 2004) it is assumed as an approximation that the year-specific excess relative radiation risk (ERR) of cancer incidence for a specified organ, defined conceptually in terms of the ratio of the relevant hazard functions, has a product form, i.e.

\[ \text{ERR} = AB. \]  

Here \( A \) depends on radiation parameters (e.g. radiation type, dose, and dose-timing), while \( B \) depends on time since irradiation and on demographic factors (e.g. age at irradiation, gender, ethnicity) but not on the dose or dose-timing. Implicit in Eq. (1) is the idea that there are really two time scales involved: \( A \) is determined by processes that occur during a comparatively short irradiation time; \( B \) by processes that occur subsequently, during a comparatively long latency period before tumors are incident clinically. An important implication of Eq. (1) is that various predicted ERR dose–response curves all have the same shape (i.e. differ only in vertical scale): one curve for dose-dependence of damage comparatively soon after irradiation; others for dose-dependence of ERR during each particular subsequent year.

“Biologically-based” radiation carcinogenesis models (e.g. Hanin et al., 2006; Heidenreich et al., 2004; Moolgavkar and Luebeck, 2003; Pierce, 2003; Sachs et al., 2005; Yakovlev and Polig, 1996) analyze the longer-time latency period in more detail. Such biologically based models usually do not assume or imply the product form, Eq. (1), for the ERR; they often do have this product form as an approximation. They can also be used when radiation is so protracted it involves times comparable to the latency time, in which case the two time scale assumption underlying Eq. (1) does not apply. However, such biologically based models are as yet less thoroughly explored and less accepted than the models actually used in current applied risk estimation, which do assume Eq. (1), as we consequently also will in the present paper.

For a single dose of magnitude \( d \) administered acutely (i.e. rapidly compared with endogenous cellular times such as DNA repair times or cell cycle times) the dose-dependent factor \( A \) in Eq. (1) has in the past usually been taken to have the following “linear-quadratic-exponential” form (reviewed in Bennett et al., 2004; Dasu et al., 2005; Radivoyevitch et al., 2001):

\[ A = (ad + bd^2) \exp(-2d - \beta d^2). \]  

Here \( a, b, \alpha \) and \( \beta \) are non-negative parameters. This form is usually rationalized, in terms of cell initiation and inactivation, as follows:

(a) Putatively, the first step in carcinogenesis by ionizing radiation is the “initiation” of target cells to form cells, that are “pre-malignant” in the sense that they may eventually evolve into a clinically detectable cancer. The molecular nature of the initiation event and the biological scenario for subsequent evolution of the initiated cells are left open in almost all applications of Eq. (2), the emphasis being instead on the numerical values of the four parameters and on the resulting dose dependence. For example, Eq. (2) does not specify whether the radiation-produced initiation is a single point mutation, much larger-scale genome damage such as aneuploidy, or some other kind of event.

(b) The factor \( \exp(-ad-\beta d^2) \) is the standard linear-quadratic (LQ) estimate of cell survival—the probability that a cell is not inactivated by the dose (i.e. is still capable of originating a clone). The quadratic term \( \beta d^2 \) represents “two-track action” where damage from two different radiation tracks interacts to inactivate the cell; the linear term \( ad \) represents one-track inactivation (reviewed in Guerrero et al., 2002; Jones et al., 2001; Sachs and Brenner, 1998; Sachs et al., 1997). In most environmental or occupational risk estimation situations, the relevant doses are so low that \( \exp(-ad-\beta d^2) \approx 1 \) to good approximation, but for second cancer scenarios such is by no means the case.

(c) The factor \( (ad+bd^2) \) is a standard LQ estimate for the product \( pN \), where \( N \) is the number of target cells and \( p \) is the probability a target cell is initiated to make a pre-malignant cell. Here \( p \ll 1 \) since \( N \), e.g. the number of stem cells in a normal breast, is believed to be \( 10^7 \) or more (Clarke, 2005; Paguirigan et al., 2006) whereas, in the situations of interest here where stochasticity is important, the number of initiated cells has order of magnitude unity.

During the last half-century, there have been many vigorous controversies about which kind of situations can be usefully approximated by Eq. (2), and about the values of the four parameters. Current disagreements about the applicability of this equation are especially heated for the initiation term \( (ad+bd^2) \) in situations where the total doses involved are much lower than those used in radiotherapy, as illustrated by strongly contradictory views of the US and French National Academies of Science (NRC, 2005; Tubiana et al., 2005). However, using, modifying, and/or generalizing Eqs. (1) and (2) is a basic starting point of almost all current applied radiation carcinogenesis risk analyses, and these two equations lead to a useful first approximation to most more sophisticated models. We shall here also start with Eqs. (1) and (2), emphasizing modifications needed in the factor \( A \), particularly at high doses where our approach differs significantly from older
approaches, to account for radiotherapy dose-fractionation and for cell proliferation during or shortly after radiotherapy. The stochastic aspects of cellular initiation, inactivation, or proliferation are emphasized. The factor $B$, whose evaluation typically involves epidemiological data, e.g. data on the Japanese atomic-bomb survivors, will be discussed only briefly.

1.2. Fractionated radiotherapy

Most external beam, fractionated, solid tumor radiotherapy regimens have the following features:

(a) Dose-fractions lasting less than 30 min are administered daily (omitting weekends). The number $K$ of dose-fractions is typically in the range 20–45.
(b) The total prescribed dose $D$ to the tumor is in the range 45–85 Gray (Gy) (1 Gy = 1 J/kg); nearby regions of the body are unavoidably also irradiated; in some proximal regions the dose is comparable to the prescribed dose.
(c) Dose-fractions are usually equal, in which case the dose $d$ for each fraction is $d = D/K$.

For fractionated radiotherapy under these conditions, the standard extension of Eq. (2), re-derived in Appendix B in order to display explicitly the assumptions involved, is

$$A = [aD + (bD^2/K)] \exp[-zD - (\beta D^2/K)].$$

Eq. (3) is an “initiation/inactivation” equation, with the LQ factor $[aD + (bD^2/K)]$ representing initiation and the LQ factor $\exp[-zD - (\beta D^2/K)]$ representing inactivation. The factors $(1/K)$ arise basically because two-track action does not occur if the two radiation tracks are separated by more than a few hours (i.e. occur in different dose-fractions), repairable damage from the first track being almost wholly repaired before the second track arrives (review: Sachs et al., 1997). Eqs. (1) and (3) have been the main formalism for estimating radiogenic second cancer risks (recent examples include Bennett et al., 2004; Dasu et al., 2005).

1.3. Weaknesses of the initiation/inactivation equation, Eq. (3)

In Eq. (3), radiation plays a dual role, initiating some normal cells but also inactivating some initiated cells. For realistic values of the parameters, the exponential (inactivation) factor $\exp[-zD - (\beta D^2/K)]$ in Eq. (3) is so small at total doses above about 15 Gy that the predicted number of second cancers is negligible—essentially, the prediction is that there is no carcinogenesis because no pre-malignant cells survive. However, recent data indicates that in fact substantial second carcinogenesis can occur at high total doses, such as those used in radiotherapy (reviewed in Sachs and Brenner, 2005; Schneider and Kaser-Hotz, 2005). A likely source of this discrepancy is that Eq. (3), as shown explicitly by its derivation (Appendix B), neglects cell proliferation between dose-fractions and during the recovery period following the last fraction. Wheldon and co-workers (e.g. Lindsay et al., 2001) pointed out that, to the contrary, repopulation by cell proliferation, which is a very well-known adverse factor for radiotherapeutic eradication of primary tumors (reviewed in McAneney and O‘Rourke, 2007), almost certainly also influences second cancer induction. Symmetric proliferation of normal and of pre-malignant stem cells is expected to increase carcinogenesis (Fig. 1).

1.4. Deterministic and stochastic initiation/inactivation/proliferation (IIP) models

To explain epidemiological data on solid second tumors, cellular repopulation by proliferation was added to the

![Fig. 1](https://example.com/f1.png)
initiation/inactivation model given by Eqs. (1) and (3). The resulting initiation/inactivation/proliferation (IIP) model (Sachs and Brenner, 2005) was based on the following equations: (a) Eq. (2) to describe initiation in each dose-fraction; (b) standard LQ equations to describe inactivation of normal and of radiation-initiated pre-malignant stem cells in each dose-fraction (Appendix B); and (c) a system of two non-linear ordinary differential equations to describe cell repopulation dynamics between dose-fractions or after the last dose-fraction (Appendix B). This model gave results consistent with data on radiation-induced breast or lung cancers following radiotherapy treatment for Hodgkin disease. However, the model neglected stochastic fluctuations of the pre-malignant cell number.

Such fluctuations may play a key role. For a related problem—tumor eradication by fractionated external-beam radiotherapy—stochastics have been studied in detail for some time (Tucker et al., 1990). An explicit analytic solution for a relevant time-inhomogeneous birth–death process with additional fractionated cell killing has been found (Zaidi and Minerbo, 2000; Hanin, 2004 and references therein). Our calculation here differs, mainly because we include initiation, but it is known (Little, 2007; Sachs and Brenner, 2005; Shuryak et al., 2006; Tucker and Taylor, 1996) that in either situation, for fractionated radiation with large total doses, repeated cycles of inactivation and proliferation can lead to statistical distributions of cell numbers where the ratio of variance to mean is much larger than the value 1 that a Poisson distribution would imply (“overdispersion”; compare Boucher et al., 1998). Correspondingly, the zero-class probability, interpreted in our setting as the fraction of patients who do not have any radiation-initiated, pre-malignant cells, can be much larger than anticipated from the mean number of radiation-initiated cells per patient. These considerations correspond to a typical eradication scenario: if, just after the last dose-fraction, every cell in every radiation-initiated pre-malignant clone has been inactivated, by radiation or other mechanisms, then subsequent cellular repopulation does not add any radiation risk.

1.5. Preview

The present paper concerns carcinogenesis in second cancer scenarios, taking into account stochastic inter-patient fluctuations in pre-malignant cell number for patients who are otherwise identical. We shall first review the deterministic initiation/inactivation/proliferation model (IIP model). Then we describe and apply a stochastic version of the model. In the stochastic model we analyze clones initiated by the radiation during a particular dose-fraction and the distribution of cell numbers for such a clone. A clone can ultimately lead to cancer incidence with some probability, with different clones presumably acting independently since they will typically originate at different random locations in an organ. In principle the way the probability of ultimate cancer incidence depends on the number of cells in a clone would need to be specified. Here we consider a limiting case which is the opposite extreme of the deterministic case—i.e. corresponds to the most pronounced stochastic effects. Specifically we assume that any clone which is present after repopulation has run its full course ultimately gives rise to a second cancer. This limiting case is assumed, explicitly or implicitly, in many biological analyses of “cancer stem cells” (reviewed in Lynch et al., 2006). It is considered here because it minimizes the number of adjustable parameters and the actual situation is expected to be intermediate between the situation predicted by the deterministic model and this limiting case stochastic model.

2. Methods

2.1. Deterministic IIP model

The deterministic IIP equations (Appendix B) deal with time-dependent average numbers, $n(t)$ and $m(t)$, of normal and initiated stem cells, respectively, for a specified organ. As will be discussed later, these equations are implied by the equations of the stochastic IIP model, and the deterministic model gives considerable insight into the stochastic one. In the models it is assumed that the excess absolute risk in any one year is much less than unity. Instead of using Eq. (3), for the deterministic model the factor $A$ in Eq. (1) is taken to be proportional to $m_{\text{final}}$, the value of radiation-induced pre-malignant cell number $m(t)$ at the “final” time, i.e. the time when the normal cell number $n(t)$ has effectively returned to its set point and post irradiation repopulation has effectively run its full course (Fig. 2). By Eq. (1) the dose-dependence of $m_{\text{final}}$ determines the shape of the predicted ERR dose-response curve. The deterministic IIP model is a special case of a somewhat more general model (Appendix A).

To calculate $m_{\text{final}}$ we previously assumed that, when $m$ is negligible compared to $n$, repopulation between dose-fractions (or during the recovery period following the last dose fraction) is described by the following differential equations (Sachs and Brenner, 2005):

\begin{equation}
\frac{dn}{dt} = F(n)n, \quad \text{where } F(n) = \lambda [1 - (n/N)],
\end{equation}

\begin{equation}
\frac{dm}{dt} = rF(n)m.
\end{equation}

In Eqs. (4) and (5) the following hold: $\lambda$ is a constant representing the maximum per-cell proliferation rate; $F(n) = \lambda [1 - (n/N)]$ is a standard logistic factor with the constant $N$ representing a set point number of normal stem cells at risk; $r$ is a constant, the “relative fitness”, describing any growth advantage or disadvantage pre-malignant cells may have compared to their normal counterparts; and in our applications $m \ll n$ at all times. The equations incorporate the idea that, with $m \ll n$, the “density” effects described by $F$ are effectively determined by the size of $n(t)$.
Because they take advantage of \( n \gg m \), our equations are quasi-linear in the following sense: the non-linear equations for normal cell number \( n(t) \) do not involve \( m(t) \), and can be solved first. Then the equations for \( m(t) \) are linear in \( m(t) \), with coefficients that depend on \( n(t) \).

2.2. Stochastic initiation/inactivation/proliferation (IIP) model

A corresponding stochastic model is given in Appendix C. For computational speed, normal stem cells are described only via their average number—the equations for \( n(t) \) are taken over without change from the deterministic IIP model. The number of initiated cells, however, is described by an integer-valued random function \( m(t) \), using the following assumptions:

(a) Inactivation by one dose-fraction of a preexisting pre-malignant cell is described by a Bernoulli distribution with parameter \( \exp(-2d-\beta d^2) \).

(b) During a dose-fraction, some cells are newly initiated and survive the fraction; these have a Poisson distribution with parameter \((ad+bd^2)(n^-/N)\exp(-ad-\beta d^2)\)

where \( n^- \) is the number of normal cells present just before the dose-fraction; here the factor \((1/N)\), which could have been absorbed in the adjustable parameters \( a \) and \( b \), is inserted for later convenience.

(c) Between dose-fractions, and after the last fraction, initiated cells undergo a time-inhomogeneous Feller–Arley birth–death process whose parameters depend on the density of normal cells; specifically, the birth rate minus the death rate at any instant is taken to be the deterministic proliferation rate, \( r/\lambda(1-[n(t)/N]) \).

(d) Standard assumptions hold on independence of the random variables involved.

(e) ERRs are calculated by assuming that the factor \( A \) is proportional to the average number of patients who have at least one radiation-produced pre-malignant cell at the “final” time defined in the caption to Fig. 2. Thus it is the presence or absence of pre-malignant cells at the “final” time, not their average number as in the deterministic IIP model, that is assumed to determine radiogenic excess risk.

As in the deterministic IIP model (Sachs and Brenner, 2005), the ERR is calculated for comparatively low doses using atomic-bomb survivor data and standard methods for “translating” the results from a Japanese to a Western cohort (Land et al., 2003). Because \( B \) is dose-independent and all our final estimates involve the product \( AB \), not either factor separately, the following steps then suffice to give ERRs at higher doses.

Using a mixture of Monte Carlo and analytic methods, we follow a clone of cells whose most recent common ancestor is a cell initiated in the \( k \)th fraction \( (k = 0, 1, \ldots, K) \); here \( k = 0 \) refers to any pre-malignant cells that
may have been present before treatment starts). Using assumptions (a) and (c) above, and iterating over the remaining dose-fractions, we calculate the probability distribution for the number of cells in such a clone at the final time. In particular this distribution gives the probability that a clone will become extinct by the final time, and gives the mean number of cells per clone at the final time. From these two quantities, we calculate, by conditioning on fraction number $k$ as discussed in detail in Appendix C, the three quantities of main interest: the mean number of pre-malignant clones per patient at the final time; the mean number of pre-malignant cells per patient at the final time (due to quasi-linearity this number turns out to be the same as the number predicted by a deterministic IIP model with the same parameters); and the “zero-class” probability, i.e. the fraction of patients who have no radiation-initiated clones in the relevant organ at the final time. This procedure gives $A$ at any dose, and calculating $AB$ at low doses from the atomic-bomb survivor data gives $B$, completing the estimate.

### 2.3. Data sets and analysis

We analyzed two data sets, chosen mainly because dosimetry information was more detailed than in other studies, in the literature on breast cancer incidence in women who had received radiotherapy for Hodgkin disease. One (Travis et al., 2003) was a matched case-control study within a cohort of 3817 female 1-year survivors of Hodgkin disease diagnosed at age 30 years or younger, between January 1, 1965, and December 31, 1994, and within 6 population-based cancer registries. Record-linkage techniques were used to identify women who developed a second primary breast cancer. For each documented case, at least two controls were selected by stratified random sampling from the cohort. Matching factors were registry, calendar year of Hodgkin disease diagnosis, age at Hodgkin disease diagnosis, and length of survival without a second cancer at least as long as the interval between the diagnoses of Hodgkin disease and breast cancer in the case. Using dose reconstruction techniques, doses were estimated both to the specific location in the breast where cancer developed for each case, and to the corresponding random variable.

The other data set (van Leeuwen et al., 2003) was a nested case-control study for a cohort of 770 female patients who had been diagnosed with Hodgkin disease before age 41 between 1965 and 1988. Detailed treatment information and data on reproductive factors were collected for 48 case patients who developed histologically confirmed breast cancer 5 or more years after diagnosis of Hodgkin disease and 175 matched control subjects. The radiation dose was estimated to the area of the breast where the case patient’s tumor had developed and to a comparable location in matched control subjects. Relative risks of breast cancer were calculated by conditional logistic regression. Follow-up as to the recent medical status of the patients was estimated to be complete for 91% of the cohort members. Six hundred and fifty of those patients survived 5 or more years.

Results of the stochastic IIP model were fitted to both data sets simultaneously. For the present, proof of principle, calculations we held all relevant parameters except $r$, $\alpha$ and $\beta$ fixed at the deterministic values; for selected values of $\alpha$ and $\beta$, $r$ was determined using a least-squares algorithm weighted with inverse estimated variance.

### 3. Results

#### 3.1. Rescaling

The IIP equations (Appendices B and C) describe inactivation and initiation by an acute dose, followed by partial repopulation, followed by another acute dose, followed by more repopulation, etc. (Figs. 1 and 2). In all essential calculations, normal cell number $n(t)$ and set point number $N$ always appear only in the ratio defining the rescaled normal cell number, $v = n/N$, never separately or in any other combination.

#### 3.2. Superposition principles for initiated cell numbers

Due to quasi-linearity as defined below Eq. (5), typical superposition results hold for the average pre-malignant cell number $m(t)$ and the corresponding random variable $m(t)$. For example, the final average number of premalignant cells is a sum of two terms. One term is due to cells which were present before treatment started and were then subject to all the cycles of inactivation and repopulation; the other term is due to cells that were initiated by one of the dose-fractions.

#### 3.3. Theorem on proliferation compensating for inactivation

In the deterministic IIP model consider the case where $r = 1$, so that normal and pre-malignant cells have identical repopulation dynamics; then the predicted dose-dependence of second cancer ERR is the same as if neither inactivation nor proliferation occurred at all. Specifically Theorem 1 in Appendix A implies that for the special case $r = 1$

$$m_{final} = K(ab + bd^2).$$

Here $m_{final}$ is the average number of radiation-induced pre-malignant cells at the “final” time (Fig. 2); and $(ad+bd^2)/N$ is the probability of initiating one normal cell
in the $k$th dose-fraction to make one pre-malignant cell just after the fraction, without regard for the fact that actually some of the newly initiated cells are also inactivated by the same dose or that some of the target cells may have been inactivated by previous dose-fractions—such inactivation is canceled out by subsequent proliferation. If $b = 0$, then $m_{\text{final}}$ is just proportional to total dose $D = Kd$. Fig. 2 shows a special case illustrating the theorem. For $r \neq 1$ however, no simple formula such as Eq. (6) for $m_{\text{final}}$ is known and presumably none exists; numerical methods are used to obtain $m_{\text{final}}$.

![Fig. 3. Different models of ERR. Four models of ERR are shown. We show the case where all four agree at low doses, i.e. the slopes of all four curves are the same at the origin, because all four models use renormalization (based on atomic bomb survivor data) at low doses. When extrapolated to higher doses, the models give different results. The parameters used are: $K = 25$ acute dose-fractions, starting on a Monday and continuing daily except for Saturdays and Sundays; initiation factors $a = 0.004 \text{ Gy}^{-1}$, $b = 0$; relative fitness $r = 1.2$; inactivation constants $\lambda = 0.1 \text{ Gy}^{-1}$ and $\beta = 0$; rate constant $c = 0.3 \text{ day}^{-1}$; and ratio of death rate to birth rate $\frac{d}{C_0} = 0.2$. Only the stochastic initiation/inactivation/proliferation (IIP) model requires all of these parameters; for example Eq. (3) contains only $a$, $b$, $\lambda$, and $\beta$. To show qualitative trends, the figure here compares different models holding common parameters fixed. If any one of the models is used in fitting data, some parameters are adjusted to fit the situation, and the adjustments would usually lead to different parameters for different models fitting the same data (see Fig. 5 for an example). Reading from top to bottom, the deterministic IIP model (blue curve), using mean pre-malignant cell number, shows an increase in ERR at high doses. This is attributed to a growth advantage that the pre-malignant cells have ($r > 1$), which comes into play especially at high doses. The linear model (dashed red line) just extrapolates the low-dose slope to high doses. According to the compensation theorem proved in Appendix A, a deterministic IIP model with $r = 1$ (instead of $r = 1.2$) and any values for its other parameters would give this linear curve. The stochastic IIP model (black solid curve), based on presence or absence of pre-malignant cells, has slope decreasing as dose increases, despite the growth advantage. Finally, the older model (dotted blue curve), given by Eq. (3), predicts almost no ERR at high doses, putatively due to inactivation of pre-malignant cells wholly uncompensated by proliferation (Fig. 1).]

3.4. Stochastic vs. deterministic results

Fig. 3 shows some representative results for the stochastic IIP model, compared to the deterministic IIP model having the same parameters and to two other models that are often used. The models differ in the way that they extrapolate lower dose estimates, based on Japanese atomic-bomb survivor data, to the higher doses also relevant in second cancer scenarios. A key point in Fig. 3 is that, even assuming initiated, pre-malignant stem cells have a growth advantage over normal stem cells during repopulation (i.e. to proliferation following inactivation), the ERR curve for the stochastic IIP model here has monotonically decreasing slope. On the other hand, when $r > 1$, the predicted curves of the deterministic IIP model always have a slope that increases as dose increases. Increasing slope contradicts epidemiological estimates, so that in the deterministic model $r \leq 1$ has previously been assumed (Sachs and Brenner, 2005).

Fig. 4 shows a reason for the difference between the deterministic and stochastic estimates. Calculations using the stochastic IIP model predict that as the dose increases, what increases is not so much the number of patients having surviving clones of initiated, pre-malignant cells but the number of pre-malignant cells per clone. This leads to situations where only a few patients have any radiation-induced pre-malignant cells, but those patients have many pre-malignant cells, corresponding to “overdispersion” in pre-malignant cell number, i.e. a variance much larger than the mean.

3.5. Modeling second cancers in Hodgkin disease patients

We previously considered data on patients treated with radiotherapy for Hodgkin disease, some of whom
subsequently developed second breast cancer (Sachs and Brenner, 2005). Fig. 5 shows that, with selected parameter choices, the stochastic IIP model can fit this data as well as the previously used deterministic IIP model. Because the stochastic model has additional adjustable parameters, the fact that it can be forced to fit the data approximately was not surprising. However, it is of interest to note that only for small values of the inactivation constants $\alpha$ and $\beta$ is an acceptable fit of the stochastic model available. For values of $\alpha$ and/or $\beta$ markedly larger than those shown in the figure caption, high doses merely lead to a few pre-malignant clones having a very large number of cells per clone. For still lower values of the radiation sensitivity to inactivation, less extreme values of the other parameters can be used in the stochastic IIP model; for example a roughly comparable fit (not shown in Fig. 5) is obtained with $\alpha = 0.04 \text{ Gy}^{-1}$, $\beta = 0$, $\lambda = 0.3 \text{ day}^{-1}$ (i.e. less rapid repopulation), $r = 1.5$, $a = 0.004 \text{ Gy}^{-1}$, $b = 0$, $c = 0.1$.

Note that the stochastic IIP model curve in Fig. 5 incorporates a growth advantage for the pre-malignant cells during the repopulation period (i.e. $r > 1$). Unless such a growth advantage is assumed, the stochastic model gives predicted high-dose values too small to match the pattern of the data.

4. Discussion

4.1. Summary

We have reviewed the deterministic IIP model and presented a stochastic version. The most important new conclusion from the stochastic IIP model is that a growth advantage for initiated and thus pre-malignant cells ($r > 1$) is compatible with ERRs that increase less rapidly than linearly at high doses (e.g. Fig. 5). In all cases thus far analyzed in sufficient detail to make parameter estimates, the deterministic IIP model gave values $r \leq 1$ (Sachs and Brenner, 2005; additional data analysis not shown). That is, during the radiotherapy and subsequent repopulation periods pre-malignant cells apparently, according to the deterministic calculations, do not have a growth advantage over their normal counterparts, whatever may happen on a longer-time scale. This result from the deterministic model was somewhat puzzling. That initiated, pre-malignant cells do have a growth advantage even on short time scales following radiation inactivation was suggested earlier (Crawford-Brown and Hofmann, 1990), found with parameter estimates using the two-stage clonal expansion model (e.g. Heidenreich, 2002), and seems plausible since on long time scales hyperplasia is a common feature of pre-malignant cells. We found that with the stochastic model this puzzling feature of the IIP model, i.e. the estimate that pre-malignant cells have no growth advantage during repopulation, is removed.

This and some other features of the stochastic model correspond to overdispersion, where the number of pre-malignant cells per patient has a variance much larger than its mean (although the number of clones per patient is Poisson-distributed, as discussed in Appendix C). This overdispersion result is consistent with previous estimates (Sachs and Brenner, 2005) and with findings of overdispersion in models, incorporating cell migration, applicable to second cancers that are leukemias (Little, 2007; Shuryak et al., 2006).

The low values of the inactivation parameters needed to bring about a fit between the stochastic IIP model and the data were a surprise. This result may point to extra radioresistance on the part of breast stem cells, as has indeed been directly observed (Phillips et al., 2006). Possibly however, the result points to the fact that the current model, where even one radiation-initiated pre-malignant cell ultimately leads to cancer, is only a limiting case where stochasticity has maximum influence. The deterministic model gives an acceptable fit for more typical values of the inactivation parameters (Sachs and Brenner, 2005) so a model intermediate between the limiting case and deterministic models would not necessarily require small values.
One interesting point emerged concerning treatment on weekends. It is often argued that, implementation difficulties apart, treating at least 6 days a week would lead to significant improvements in tumor control probabilities. Our analysis shows that weekend treatment gaps likewise adversely affect the risk of second cancers: repopulation during weekends tends to increase the number of pre-malignant cells right after the end of the last dose-fraction, which is the key time for extinction.

### 4.2. Some weaknesses of the IIP models

The models presented here, and more specifically the stochastic IIP model, have various weaknesses, including the following:

(a) The product estimate for ERR, Eq. (1), is only a phenomenological way to model tumor progression during the comparatively long latency period which follows irradiation.

(b) Even given the product assumption, taking the corresponding dose-dependent factor $A$ in the stochastic model to be, in effect, a binary variable, with value zero or one according as a patient has no pre-malignant cells or any number of pre-malignant cells, respectively, gives only the limiting case where stochastic effects are maximal.

(c) The effects of radiation on pre-existing pre-malignant cells are neglected in the analysis. For high ERRs at doses high enough for significant cell killing per dose fraction, as in the data of Fig. 5, this is a reasonable approximation; for situations where the sensitivity to radiation-induced cancer is less, the effects would have to be taken into account and, in the stochastic IIP model, would lower the ERR prediction somewhat.

(d) Intercellular interactions are taken into account only in one way, via a single logistic factor. The actual richness of intercellular signaling and cellular reaction to microenvironments is not considered.

(e) No molecular mechanisms are modeled. As far as the IIP models are concerned, a cell might as well be a very simple object capable only of proliferation, being initiated, and being inactivated. The models in their present form work equally well (or equally badly) whether initiation is interpreted as a single point mutation or as any other somatically heritable change. With minor alterations the models could be applied even if initiation involves triggering of a multi-cellular reaction such as angiogenic recruitment to a dormant tumor.

(f) For solid tumors, no spatial properties are taken into account. Effects of dose-inhomogeneity can be taken into account with dose-volume histograms (Koh et al., 2007) but spatial factors during tumor progression are more complicated (e.g. Enderling et al., 2007).

(g) Effects of intrinsic inter-patient heterogeneity are not taken into account.

(h) Many of these weaknesses were previously accepted in the interests of keeping the number of adjustable parameters so small that genuine predictions are possible (Sachs and Brenner, 2005). However, the stochastic IIP involves additional adjustable parameters, so many that it is not presently possible to determine them all by other data and thereby allow clear predictions when dealing with second cancers.

### 4.3. Conclusions

Second cancers after radiotherapy are of increasing concern. They are influenced by cellular repopulation during and shortly after treatment. The IIP models are the first systematic, quantitative approach based on cell population dynamics including repopulation for realistically estimating second cancer risk after fractionated irradiation. Consequently, incrementally improving the IIP models will be worthwhile. A deeper understanding of the initiation/inactivation/proliferation process that apparently underlies radiation-induction of solid tumors at high doses should lead to additional insights, potentially suggesting practical improvements in radiotherapy.

Such investigations should also clarify fundamental carcinogenesis processes in humans. Modeling second cancers has an important advantage as regards increasing our basic understanding, compared to analyzing animal, in vitro, or in silico data: one deals directly with the endpoint of main interest, human cancer, not surrogate endpoints requiring difficult extrapolations. Especially important in improving biologically based second cancer models will be data on intermediate endpoints such as the number and size of hyperplastic foci during the years after radiotherapy. Whether and how modern high-throughput molecular data can be used remains to be seen.

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### Appendix A. General deterministic formalism

This appendix first presents a deterministic formalism that generalizes our previous formalism (Sachs and Brenner, 2005); then we correspondingly generalize a previous theorem on cellular proliferation compensating for cellular inactivation.
A.1. Equations

The general formalism differs as follows from the deterministic model used previously (Sachs and Brenner, 2005):

(a) Time between fractions is adjustable so that irregularly timed dosing can be included, and similarly the dose per dose-fraction is allowed to vary (e.g. extra dose given during certain dose-fractions).

(b) Radiation cell inactivation for one dose-fraction need not be LQ, but rather can involve any non-linearities, e.g. the possible low-dose non-linearities now being intensively investigated (Hall, 2004).

(c) Radiation cell initiation in one dose-fraction can depend on the dose in that fraction in any way, not just via a linear-quadratic-exponential expression as in Eq. (2).

(d) Repopulation, between doses and after the last dose, can follow any restorative pattern, not just the logistic pattern of Eq. (4) or similarly specific patterns that are often postulated (e.g. Gompertzian as in Wheldon et al., 2000).

(e) The assumption \( m \ll n \), made as a separate assumption in the earlier paper, is incorporated into the basic equations ab initio.

Thus we consider a population of \( n(t) \) normal cells and \( m(t) \) initiated cells in an organ receiving clinically significant doses during fractionated external-beam radiotherapy. We have in mind the interpretation that \( n \) refers to stem cells and \( m \) to “pre-malignant stem cells”. Denote the time of the \( k \)th dose-fraction by \( t(k) \) and the dose by \( d(k) \). We assume that:

(a) After the \( k \)th fractionated dose, the surviving fraction \( S(k) \) for preexisting normal and pre-malignant cells is the same, i.e. \( S(k)n^-(k) \), respectively, \( S(k)m^-(k) \), survive the dose; here \( n^-(k) \), respectively, \( m^-(k) \), is the number present just before the \( k \)th dose-fraction and \( 0 \leq S(k) \leq 1 \). The dependence of surviving fraction \( S(k) \) on dose \( d(k) \) per dose-fraction remains unspecified in this general model; one could have a typical LQ surviving fraction \( S(k) = \exp[-ad(k) - \beta d(k)] \) as in Eq. (2) of the text, or have some more complicated dose dependence.

(b) Between doses and after the last dose, the per-cell repopulation rate of pre-malignant cells, corresponding to symmetric division (compare Shuryak et al., 2006), is a constant, \( r \), times the per-cell repopulation rate for normal cells. Here repopulation rates refer to repopulation during comparatively short time periods (e.g. a day or a weekend between doses, and a number of weeks after the last dose). Growth during the much longer latency periods involved in the development of clinical cancer from pre-malignant cells could in general have different dynamics.

(c) Because typical situations involve a normal cell number \( > 10^6 \) and a pre-malignant cell number \( < 10^3 \), we assume that \( m(t) \ll n(t) \) throughout, and that the fraction of normal cells that is initiated to produce pre-malignant cells is so small it can be neglected compared to the total number of normal cells. This assumption allows us to track the time-evolution of normal cell number independently of the time-evolution of initiated cell number (though not vice versa).

(d) Cellular migration, important in leukemogenesis after partial body high-dose radiation (Shuryak et al., 2006) but much less important for solid tumors, is neglected.

The idea underlying assumptions (a) and (b) is that the pre-malignant cell population is derived from the cell population at risk by an initiating event which need not affect radiation survival markedly, or drastically affect the growth characteristics during comparatively short periods of repopulation in response to cell killing.

Let \( n^+(k) \) denote the number of normal cells just after the \( k \)th fractionated dose. Then by assumption (a) above:

\[
 n^+(k) = S(k)n^-(k), \quad 1 \leq k \leq K. \tag{A.1}
\]

Here we neglect the decrease of \( n^+(k) \) due to the very small fraction of at-risk cells that is initiated by the radiation (assumption (c)). The range \( 1 \leq k \leq K \) will apply throughout, unless explicitly stated to the contrary.

The number \( m^+(k) \) of altered cells just after the \( k \)th dose depends, by our assumption (a), on the same factor \( S(k) \):\n
\[
 m^+(k) = S(k)m^-(k) + T(k)S(k)n^-(k). \tag{A.2}
\]

Here \( T(k) \), with \( T(k) \ll 1 \), denotes the fraction of at-risk cells that are initiated, so \( T(k)S(k) \) is the fraction that are initiated and also survive the dose-fraction; the condition \( T(k) \ll 1 \) corresponds to our blanket assumption \( m \ll n \). The dependence of \( T(k) \) on dose \( d(k) \) for the \( k \)th dose-fraction need not be specified. For example, for a given normal number setpoint \( N \), \( T(k) \) could have the LQ form

\[
 T(k) = 1 - \exp\left[-[ad(k) + b d^2(k)]/N\right]
\]

in which case, since \( T(k) \ll 1 \), \( T(k) \approx [ad(k) + b d^2(k)]/N \) and in this approximation \( NT(k)S(k) \) has the form given in Eq. (2) of the main text if \( S(k) \) is LQ. Or the form of the initiation factor \( T(k) \) could be more general.

Repopulation of normal cells between dose-fractions and for several weeks after the last fraction will be modeled using a per-cell repopulation rate \( F \) generalizing the logistic form \( F = \lambda[1 - (n(t)/N)] \) discussed in connection with Eq. (3) of the main text. Thus we shall assume

\[
 \frac{dn}{dt} = F(n)n \Rightarrow n^+(k+1) = R(k)n^+(k), \quad \text{where } R(k) = \exp\left[\int_{t(k)}^{t(k+1)} F(n(t))dt\right]. \tag{A.3}
\]

Here \( t(k) \) and \( t(k+1) \) are the respective times of the \( k \)th and the \((k+1)\)th fractions; if \( k = K \) then \( t(k+1) \) is taken as \( t(K+1) = \infty \), interpreted as a time some weeks after the final dose-fraction when repopulation has
effectively run its full course (compare Fig. 2) and the slower phase of carcinogenesis, modeled in this paper only by the factor R in Eq. (1), begins. If F has the prototype logistic form \( F = \frac{1}{2} [1-(n/N)] \) integration gives for \( R(k) \) in Eq. (A.3), \( R(k) = N/[n^+(k)[1-x] + Nx] \), where \( x = \exp[-\lambda(k+1)-\tau(k)] \); then for \( t(k+1) \to \infty \), \( x \to 0 \) so \( R \\to N \). Thus in this case \( R(k) \) is a repopulation factor obeying:

\[
\begin{align*}
\text{(a)} & \quad \text{if } n^+(k) < N, R(k)>1, \\
\text{(b)} & \quad \text{for } t(k+1) \to \infty, \quad R(k)n^+(k) \to N. 
\end{align*}
\]

(A.4)

Generalizing to include many other reasonable growth patterns (such as Gompertzian), we will leave \( F \) general for the time being but assume throughout that for \( R(k) \) as defined in Eq. (A.3), condition (b) in Eq. (A.4) holds. The general deterministic model is completed by using Eq. (5) of the main text verbatim, i.e.

\[
dn/dt = rF(n)n. 
\]

(A.5)

We showed earlier (Sachs and Brenner, 2005) that manipulating Eq. (A.5) and the differential equation \( dn/\text{d}t = F(n)n \) in Eq. (A.3) implies a simple relation between the way normal and pre-malignant cells repopulate, namely:

\[
m^−(k + 1) = m^+(k)[n^−(k + 1)/n^+(k)]^r. 
\]

(A.6)

A.2. General deterministic model: compensation theorem

We now generalize a previous theorem (Sachs and Brenner, 2005). For generality we include pre-malignant cells that may have been present before the start of radiotherapy. In view of the fact that background carcinogenesis and radiation carcinogenesis produce the same spectrum of cancer types (Little, 2000) we treat such pre-existing pre-malignant cells on the same footing as radiation-induced pre-malignant cells. The theorem states that if \( r = 1 \) then, by the time repopulation has run its full course, repopulation has completely compensated for cell inactivation as far as the number of pre-malignant cells is concerned (Fig. 1). More formally, we have the following:

**Theorem 1.** Suppose Eqs. (A.1)–(A.5) hold and \( r = 1 \). Let \( m_0 = m^−(1) \) be the number of pre-malignant cells just before therapy starts. Then

\[
m(\infty) = m_0 + N \sum_{k=1}^{K} T(k) 
\]

(A.7)

dependent on the initiation factors \( T(k) \), but independent of the inactivation factors \( S(k) \) and the repopulation factors \( R(k) \), and thus equal to the result of a hypothetical process where neither inactivation nor repopulation occurs.

The proof consists of iterating Eqs. (A.1)–(A.5) to get the time course for \( m \) and \( n \). Just before the first dose-fraction \( n \) has its set point value, i.e. \( n^−(1) = N \). Just afterwards we therefore have:

\[
n^+(1) = S(1)N, \quad m^+(1) = S(1)[m_0 + T(1)N]. 
\]

(A.8)

Eq. (A.8) and \( r = 1 \) in Eq. (A.6) show that just before the second fraction

\[
n^−(2) = R(1)S(1)N, \quad m^−(2) = R(1)S(1)[m_0 + T(1)N]. 
\]

(A.9)

Eq. (A.9) in turn gives two key results: \( m^+(2) = S(2)R(1)S(1)N \); and

\[
m^+(2) = S(2)R(1)S(1)[m_0 + T(1)N] + T(2)S(2)R(1)S(1)N 
\]

\[
= S(2)R(1)S(1)[m_0 + (T(1) + T(2))N]. 
\]

(A.10)

The crux of the entire argument is the fact that the term \( T(2)N \) (which refers to initiation by the second dose-fraction) and the term \( m_0 \) (which refers to pre-malignant cells present prior to therapy), are both multiplied by same factor, \( S(2)R(1)S(1) \) as is the term \( T(1)N \). This result was initially somewhat surprising to us, since the factor \( R(1)S(1) \) refers to inactivation by the first dose-fraction and subsequent repopulation, which seem at first blush to have no relation to initiation by the second dose-fraction; there is an indirect relation because initiation during the second dose-fraction is proportional to the number of normal cells present just before that fraction, which is influenced by inactivation during the first dose-fraction and subsequent repopulation.

By a simple induction argument we now get

\[
\begin{align*}
\text{(a)} & \quad n^+(K) = \Pi N, \\
\text{(b)} & \quad m^+(K) = \Pi \left[ m_0 + N \sum_{k=1}^{K} T(k) \right], \\
\text{where } & \quad \Pi = \prod_{k=1}^{K} S(k) \prod_{k=1}^{N-k} R(k). 
\end{align*}
\]

(A.11)

Combining \( \Pi = N, \) Eq. (A.6) with \( r = 1 \), and Eq. (A.11) gives

\[
m(\infty) = m^+(K)[n(\infty)/n^+(K)] = m_0 + N \sum_{k=1}^{K} T(k). 
\]

In this last relation all the factors involving killing and repopulation have contrived to cancel out, and the result is Eq. (A.7), as was to be shown. Eq. (A.7) implies that \( m_0 \) reemerges (Phoenix-like) at the final time despite many intermediate inactivation/proliferation vicissitudes.

A simple example of the theorem in a special case is shown graphically in Fig. 2.

Appendix B. The deterministic initiation/inactivation/ proliferation (IIP) model

B.1. Specializations of Appendix A

For \( m \ll n \), as holds throughout the present analysis, the formalism used previously (Sachs and Brenner, 2005), is a special case of Eqs. (A.1)–(A.5). The relevant specializations are the following:

\[
\begin{align*}
\text{(a)} & \quad \text{all dose-fractions are equal so that, for all } k, \quad d(k) = (D/K) \text{ (total dose divided by fraction number); we then write } d(k) = d. \\
\text{(b)} & \quad \text{inactivation is } LQ, \text{ so that } S(k) = \exp[-\alpha d - \beta d^2]; 
\end{align*}
\]
(c) initiation is LQ so that $T(k) = (1/N)(ad + bD^2)$, where $a$ and $b$ are non-negative adjustable constants and the factor $(1/N)$ has been inserted for later convenience;
(d) normal cell proliferation is logistic, so that $F = \lambda [1 - (n/N)]$, implying $R(k) = 1/[(n^+(k)/N)[1 - x] + x]$, where $x = \exp \{-\lambda [t(k + 1) - t(k)]\}.

B.2. Rescaling and equations

When we substitute specializations (a)–(d) into Eqs. (A.1)–(A.5) and use the indicated initial condition B.2. Rescaling and equations may assume Eqs. (A.3) and (A.5); Eq. (A.3) with in a way that does not involve $k$ normal cell number. Specifically, substituting specializations (a)–(d) gives the following results. The effect of the $k$th dose fraction on rescaled normal cell number $v(t)$ and on pre-malignant cell number $m(t)$ are given by

\[ v^+(k) = S v^-(k) \text{ with } S = \exp[-ad - \beta d^2], \]
\[ v \equiv n/N, \quad k = 1, \ldots, K, v^+(1) = 1, \]
\[ m^+(k) = S m^-(k) + [ad + bD^2]Sv^-(k). \]  (B.2)

Between dose-fractions and for several weeks after the last dose-fraction:
\[
\frac{dn}{dt} = \lambda (1 - v) \Rightarrow v^+(k + 1) = 1/[v^+(k)[1 - x] + x],
\]
\[
\text{where } x = \exp[-\lambda [t(k + 1) - t(k)]], \]
\[
\frac{dm}{dt} = r(1 - v)m \Rightarrow m^+(k + 1) = m^+(k)[v^-(k + 1)/n^+(k)]^r. \]  (B.4)

Here, for $k = K$, $t(k+1)$ again refers to the final time (Fig. 2). Quasi-linearity here shows up via the fact that Eqs. (B.2) and (B.4) contain $m(t)$ linearly. We will refer to Eqs. (B.1)–(B.4) as the deterministic initiation/inactivation/proliferation (IIP) model. Thus Eqs. (A.1)–(A.5) will be referred to as a generalization of the deterministic IIP model.

B.3. Derivation of Eq. (3)

Eq. (3) of the main text is usually derived from Eq. (2) by assuming that no repopulation occurs. In fact, setting repopulation to zero in the deterministic IIP model does imply Eq. (3), as follows. We can set repopulation to zero by putting $\lambda = 0$ in Eqs. (B.3) and (B.4). Then, $F = 0$ in Eqs. (A.3) and (A.5); Eq. (A.3) with $F = 0$ implies that $R(k) = 1$ for all $k$; and Eq. (A.5) with $F = 0$ implies that we may assume $r = 1$ without essential loss of generality. Consequently we can use Eq. (A.11), which was based on $r = 1$. Substituting into Eq. (A.11) $R(k) = 1$, $D = d/K$ from specialization (a) above, $S(k) = \exp[-ad - \beta d^2]$ from specialization (b), and $T(k) = [ad + bD^2]/N$ from specialization (c) gives:

\[ m(\infty) = m^+(K) = [m_0 + aD + (bD^2)/K] \times \exp[-aD - \beta (D^2/K)]. \]  (B.5)

Thus if $m_0 = 0$, Eq. (3) follows. Realistically speaking, however, the no-proliferation assumption $F = 0$ is not expected to hold, and using Eq. (3) is expected to give inaccurate results at high doses (compare Fig. 5).

Appendix C. The stochastic IIP model

Customized Fortran programs were used to implement the following assumptions and equations defining the stochastic IIP model, which extends the deterministic IIP model of Appendix B.

C.1. Equations for cell numbers

For the average number of normal cells and for the non-negative integer-valued random function $m(t)$ describing initiated cell number we make assumptions corresponding to the deterministic IIP model:

(a) Normal stem cell number is expected to be far greater than 1, so in our stochastic model we still analyze this number deterministically, using Eqs. (B.1) and (B.3) for the rescaled normal cell number $v$.
(b) Corresponding to Eq. (B.2) it is assumed for $m(t)$ that at the $k$th dose fraction:
(i) each cell present before the fraction has probability $1 - S$ of being inactivated by the fraction, independently of the other cells, where again $S = \exp[-ad - \beta d^2]$;
(ii) the probability of producing new, live, pre-malignant cells by initiation during that dose-fraction is given by a Poisson distribution with average $(ad + bD^2)v^-(k)S$.
(c) Between fractions and after the last fraction $m(t)$ is assumed to undergo a Feller–Arley time-inhomogeneous birth–death process. Such processes have been reviewed, e.g. by Tan (2002). They were applied to a related problem, the problem of tumor eradication by fractionated radiation, by Hanin and coworkers (Hanin, 2004; Hanin et al., 2006), who obtained exact solutions of the relevant stochastic equations in that case. We here assume the per-cell birth rate $\rho(t)$ and death rate $\delta(t)$ are given in terms of an adjustable parameter $c$ with $0 \leq c < 1$ by

\[ \delta(t) = c\rho(t), \quad \rho(t) - \delta(t) = \lambda r[1 - v(t)]. \]  (C.1)

For example, if $c$ is increased the death rate increases, but the difference between birth and death rates remains the same as in the deterministic model.

C.2. Calculating statistics for pre-malignant cells

The time dependence of $m(t)$ between dose-fractions and after the last fraction is then determined by probability
distributions involving the following time integrals:

\[
\zeta(t) = \exp \left\{ - \int_{t(k)}^{t(k+1)} \mathrm{d}r \left[ \rho(t) - \delta(t) \right] \right\},
\]

\[
\zeta_k = \int_{t(k)}^{t(k+1)} \mathrm{d}t \rho(t) \zeta(t).
\]  

(C.2)

Specifically (Tan, 2002, pp. 169–171), the probability that a clone founded by a pre-malignant cell present just after the \(k\)th dose-fraction will become extinct before the next fraction is

\[
1 - (\zeta_k + \zeta_k^{-1})^{-1}, \quad \text{where} \quad \zeta_k = \zeta(t(k+1))
\]  

(C.3)

and the probability this clone contains exactly \(j\) cells just prior to the next fraction \((j = 1, 2, \ldots)\) is

\[
\zeta_k^{j-1}(\zeta_k + \zeta_k^{-1})^{-1}.
\]  

(C.4)

Here, as before, \(t(k+1)\) for the last dose-fraction \((k = K)\) is taken formally as infinite and interpreted as a “final time” of roughly 60 days, as described in the caption of Fig. 2. Using Eqs. (C.3) and (C.4) enabled us to avoid splitting the time between dose-fractions or after the last dose-fraction into small steps, with Monte-Carlo calculations at each step; instead we needed just one Monte-Carlo evaluation for each dose-fraction in any one sample run.

To increase computational speed, algorithms were designed to minimize the number of Monte-Carlo steps. Suppose that just after the \(k\)th fraction there is exactly one pre-malignant cell. Here we allow \(k = 0\), referring to pre-malignant cells present just before treatment starts. The pre-malignant cell can give rise to a clone. The clone could grow by proliferation, or it could die out due to radiation inactivation and/or processes reflected in the death rate \(\delta(t)\), such as apoptosis. Eqs. (C.1)–(C.4), together with Monte-Carlo calculations for the number of initiated cells inactivated in each subsequent fraction, allow one to calculate numerically for the clone the following quantities:

(a) The probability distribution for the number of cells at the final time \((t = \infty)\), interpreted as several weeks after therapy starts; compare Fig. 2 and its caption for a discussion of short and long time scales and the “final time”). This probability distribution in turn determines the following two quantities.

(b) The probability \(e_k\) that such a clone becomes extinct before the final time. For non-zero death rate in the Feller–Arley process \(e_k\) is different from zero even for \(k = K\), i.e. some clones can “accidentally” die out even after radiation stops.

(c) The average number \(f_k\) of cells in a non-extinct clone at time \(t = \infty\).

These quantities, \(e_k\) and \(f_k\), in turn enable us to calculate the quantities discussed in the main text: the average number of clones per patient; and the zero-class probability, i.e. the probability a patient has no pre-malignant cells at the final time. The results are the following.

By the definition of \(e_k\), the average number of clones per patient, which we will denote by \(\text{clones}\), is

\[
\text{clones} = \sum_{k=0}^{K} y(1 - e_k), \quad \text{where}
\]

\[
y = (ad + bd^2) \exp(-zd - \beta d^2).
\]  

(C.5)

The probability that all clones initiated in the \(k\)th fraction have become extinct is the following, using our Poisson assumption on initiation, conditioning on the number initiated, and assuming the number of pre-malignant cells present prior to the start of treatment is also Poisson-distributed:

\[
\exp(-y) \sum_{l=0}^{\infty} (ye_k)^l / l! = \exp(-y(1 - e_k)).
\]  

(C.6)

The zero-class probability, which we will denote by \(\text{zero}\), is the product of these probabilities, i.e.

\[
\text{zero} = \exp \left\{ - \sum_{k=0}^{K} y(1 - e_k) \right\}.
\]  

(C.7)

We took \(A\) in Eq. (1) proportional to \(1 - \text{zero}\), thereby obtaining the dose-dependence of \(A\) apart from an overall scale factor.

Eq. (C.7) can also be derived from the more general observation that the number of surviving clones is Poisson-distributed, proved as follows. The number of clones initiated by the \(k\)th dose is, by assumption, Poisson-distributed. The number that are not eradicated by the final time is a random thinning of the number initiated, so it is also Poisson-distributed. The total number of clones initiated and surviving is thus a sum (from \(k = 0\) to \(K\)) of independent Poisson random variables, and is therefore itself a Poisson random variable. Consequently the probability of this random variable being zero is the exponential of the negative mean, i.e. \(\text{zero} = \exp(-\text{clones})\), which, by Eq. (C.5) and the fact that a patient is free of pre-malignant cells at the final time iff the patient is free of surviving pre-malignant clones at the final time, implies Eq. (C.7), as was to be shown.

Quasi-linearity implies that the average number of pre-malignant cells per patient is the same as in the deterministic theory. This implication provided an internal check on our computer algorithms.

References


