

REVIEW

Modeling and measurement of signaling outcomes affecting decision making in noisy intracellular networks using machine learning methods

Mustafa Ozen ¹, Tomasz Lipniacki², Andre Levchenko³, Effat S. Emamian⁴, and Ali Abdi^{1,5,*}

¹Center for Wireless Information Processing, Department of Electrical and Computer Engineering, New Jersey Institute of Technology, 323 King Blvd, Newark, NJ 07102, USA, ²Institute of Fundamental Technological Research, Polish Academy of Sciences, Pawinskiego 5B, 02-106 Warsaw, Poland, ³Yale Systems Biology Institute and Department of Biomedical Engineering, Yale University, New Haven, CT 06520, USA, ⁴Advanced Technologies for Novel Therapeutics, Enterprise Development Center, New Jersey Institute of Technology, 211 Warren St., Newark, NJ 07103, USA, and ⁵Department of Biological Sciences, New Jersey Institute of Technology, 323 King Blvd, Newark, NJ 07102, USA

*Corresponding author. E-mail: ali.abdi@njit.edu

Abstract

Characterization of decision-making in cells in response to received signals is of importance for understanding how cell fate is determined. The problem becomes multi-faceted and complex when we consider cellular heterogeneity and dynamics of biochemical processes. In this paper, we present a unified set of decision-theoretic, machine learning and statistical signal processing methods and metrics to model the precision of signaling decisions, in the presence of uncertainty, using single cell data. First, we introduce erroneous decisions that may result from signaling processes and identify false alarms and miss events associated with such decisions. Then, we present an optimal decision strategy which minimizes the total decision error probability. Additionally, we demonstrate how graphing receiver operating characteristic curves conveniently reveals the trade-off between false alarm and miss probabilities associated with different cell responses. Furthermore, we extend the introduced framework to incorporate the dynamics of biochemical processes and reactions in a cell, using multi-time point measurements and multi-dimensional outcome analysis and decision-making algorithms. The introduced multivariate signaling outcome modeling framework can be used to analyze several molecular species measured at the same or different time instants. We also show how the developed binary outcome analysis and decision-making approach can be extended to more than two possible outcomes. As an example and to show how the introduced methods can be used in practice, we apply them to single cell data of PTEN, an important intracellular regulatory molecule in a p53 system, in wild-type and abnormal cells. The unified signaling outcome modeling framework presented here can be applied to various organisms ranging from viruses, bacteria, yeast and lower metazoans to more complex organisms such as mammalian cells. Ultimately, this signaling outcome modeling approach can be utilized to better understand the transition from physiological to pathological conditions such as inflammation, various cancers and autoimmune diseases.

Key words: Cell decision making; noise; decision theory; machine learning; signaling errors; p53 system

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INSIGHT BOX

Cells are supposed to make correct decisions, i.e. respond properly to various signals and initiate certain cellular functions, based on the signals they receive from the surrounding environment. Due to signal transduction noise, signaling malfunctions or other factors, cells may respond differently to the same input signals, which may result in incorrect cell decisions. Modeling and quantification of decision-making processes and signaling outcomes in cells have emerged as important research areas in recent years. Here we present univariate and multivariate data-driven statistical models and machine learning methods for analyzing dynamic decision-making processes and signaling outcomes. Furthermore, we exemplify the methods using single cell data generated by a p53 system, in wild-type and abnormal cells.

INTRODUCTION

Understanding how cells make decisions in response to input signals is an important challenge in molecular and cell biology. Depending on the signals they receive, cells can adopt different fates. Emergence of single cell data and methods [1–3] has made it possible to study and model the behavior of each cell individually. An important factor that affects cell decisions is biological noise in various organisms [4], which can cause cells to exhibit different behaviors, when receiving the same input signal. For example, under the same stimuli, some cells may decide to survive, whereas others may undergo apoptosis. Signaling outcomes can be affected by genetic and epigenetic regulation and misregulation, leading to errors in signaling outcomes and ensuing cell decisions.

Given the probabilistic nature of cellular decisions [1, 3], it is of interest to have a unified set of statistical metrics and methods to systematically study and characterize the signaling outcomes that may inform them and determine probabilities associated with different outcomes. Using statistical signal processing and decision theory concepts, recently a framework was introduced by Habibi *et al.* [1], to compute optimal decision thresholds and probabilities for incorrect cell decisions using single cell data. More specifically, in the transcription factor nuclear factor κ B (NF- κ B) pathway regulated by the tumor necrosis factor (TNF) [3], the optimal decision threshold which minimized the decision probability to distinguish between two different TNF levels was computed from data [1]. Probabilities of incorrect cell decisions were computed from data as well.

One goal of this paper is to show how the statistical decision theoretic framework [1] can be used to study other molecular systems and signaling outcomes. The other goal is to extend the decision modeling framework such that one can model and analyze multi-dimensional signaling outcome processes using multi-time point measurements. This allows to incorporate signaling dynamics into decision-making analysis. Application of receiver operating characteristic (ROC) curve as a graphical tool to visualize decisions and outcomes under normal and abnormal conditions is introduced here as well. In this paper, we use the tumor suppressor p53 system, as an example, to present the concepts, metrics and algorithms related to decision-making and outcome analysis.

The tumor suppressor p53 is an important transcription factor that is responsible for DNA repair, cell cycle suppression, cell growth control and initiation of apoptosis [5–8]. When a healthy cell is exposed to ionizing radiation (IR), DNA damage occurs [9]. Due to the DNA damage, p53 becomes activated [5, 10, 11], and the cell takes one of two possible actions: it can either survive by repairing the DNA or trigger apoptosis [8, 12,

13]. Our focus here is to demonstrate how such outcomes can be systematically modeled. We accomplish this by introducing metrics and methods to evaluate success and failure rates of the signaling outcomes and actions, in response to the DNA damage caused by different IR doses and under various conditions. In order to do this, we collected data using the simulator of Hat *et al.* [9], to obtain single cell data of healthy cells, when different IR doses are applied. Moreover, we collected single cell data of abnormal cells exposed to different IR doses, to measure how the decision-making is affected when there is an anomaly in the system, in addition to the DNA damage.

The rest of the paper is organized as follows. First, we briefly explain the p53 system and its response to DNA damage. Then we present decision-making and outcome analysis as a hypothesis testing problem on the IR level, define probabilities associated with various decisions, introduce the optimal decision-maker and describe the single cell data used to determine the decision probabilities in the presence of noise and under normal and abnormal conditions. Methods for computing optimal decision thresholds and the associated decision error rates are presented afterwards, using either single, double or multiple time point measurements in individual cells. The latter is particularly useful to understand the effect of temporal variations and dynamical changes. Additionally, ROC curves are computed and presented as useful tools to visualize the trade-off between decision error rates and how they are affected by decision thresholds and other factors. A comparison between binary and ternary decision-making and outcome analysis and their error rates is provided as well. The paper concludes with a summary of the highlights of the methods and their biological implications for understanding signaling outcomes and decisions in the exemplary p53 system and extensions to other systems.

SIGNALING OUTCOMES AND DECISIONS IN THE P53 SYSTEM WHEN DNA DAMAGE OCCURS: A CASE STUDY

The transcription factor p53 has a significant role in DNA repair, cell cycle suppression, regulation of cell growth and initiation of apoptosis [5–8]. It becomes active in response to DNA damage that may occur when the cell is exposed to IR, ultraviolet radiation, heat shock, etc. [5, 10, 11]. In particular, exposure to IR results in DNA double-strand breaks (DSBs), which are the most serious DNA lesion. When DSB is not repaired, it can cause cell death or DNA mutations which can propagate to new cell generations [12, 14, 15]. When DNA damage occurs, p53 can assume two phosphorylation states: p53_{Arrester} and p53_{Killer}. Afterwards, the p53 system can take two actions: it either suppresses cell

cycle until DNA is repaired, if the damage is low and repair is possible, or can trigger apoptosis if the damage is high and repair is not possible [8, 12, 13]. Herein, we intend to compute decision thresholds and incorrect decision rates when the DNA damages caused by various IR doses occur in a cell. With this goal in mind, we conduct stochastic simulations of cells exposed to different IR doses [9], to obtain *in silico* single cell data.

Consider the p53 system model [9] shown in Fig. 1. The p53 system is activated due to a DNA damage induced by IR. Initially the protein kinase ataxia-telangiectasia mutated (ATM) is activated by the DNA damage [16, 17]. The active ATM phosphorylates Mdm2, which is a p53 inhibitor [18]. The ATM also activates p53 by phosphorylating it to one of its active phosphoforms, p53_{Arrester}, which further phosphorylates p53 to the p53_{Killer} form [19–21]. Moreover, the p53_{Arrester} activates the Mdm2 [22] and wild-type p53-induced phosphatase 1 (Wip1) [23, 24]. The active Wip1 inhibits the ATM [25] and dephosphorylates the p53_{Killer} to the p53_{Arrester} form [26]. The p53_{Killer} regulates another phosphatase, phosphatase and tensin homolog (PTEN), which initiates a slow positive feedback loop stabilizing the level of p53 [27]. If DNA damage is large and its repair takes a longer time, PTEN accumulates to high levels and inhibits AKT, which may no longer phosphorylate Mdm2. Unphosphorylated Mdm2 remains in cytoplasm and may not target nuclear p53 for degradation. Thus, accumulation of PTEN results in disconnection of negative feedback loop between p53 and Mdm2. The slow positive feedback loop acts as a clock, giving cells time to repair DNA and initiating apoptosis if DNA repair takes too long. The apoptotic module, where transcription of pro-apoptotic proteins is induced, is controlled by p53_{Killer} and Akt that suppresses the apoptosis. When Akt is inhibited by increased level of PTEN, it will no longer suppress the apoptotic module. Thus, the p53_{Killer} will initiate activation of cysteine-aspartic proteases (caspases), enzymes having an essential role in cell death (Fig. 1). Since we are interested in the analysis of the signaling outcomes which affect whether the cell survives or triggers apoptosis, we do not consider the cell cycle arrest module (regulated by p53_{Arrester}) and focus on the apoptotic module. Simulation files can be found in Hat et al. [9], and more detailed information about the p53 system and each component and interaction there can be found in Hat et al. [9] and Bogdal et al. [29]. More specifically, interested readers can refer to the Supporting Information S1 Text of [9], which includes a summary of mathematical models of the p53 system, a detailed description of the model, a notation guide and lists of parameters and reactions.

DECISION-MAKING AND OUTCOME ANALYSIS: HYPOTHESIS TESTING ON INPUT SIGNALS AND OPTIMAL DECISIONS WITH MINIMUM ERRORS

When cells are exposed to radiation, each cell may respond differently due to noise or some other factors. One may decide to survive, whereas another may trigger apoptosis, both under the same IR dose. Given the probabilistic nature of such decisions [1], we can formulate p53-based decision-making as a binary hypothesis testing problem, where the decision-making system is going to test which of the following two hypotheses is true regarding the applied IR dose, to trigger an action accordingly:

$$\begin{aligned} H_0 &: \text{IR dose is low,} \\ H_1 &: \text{IR dose is high.} \end{aligned} \quad (1)$$

Binary hypothesis testing is observed in other systems, e.g. the TNF/NF- κ B system [1].

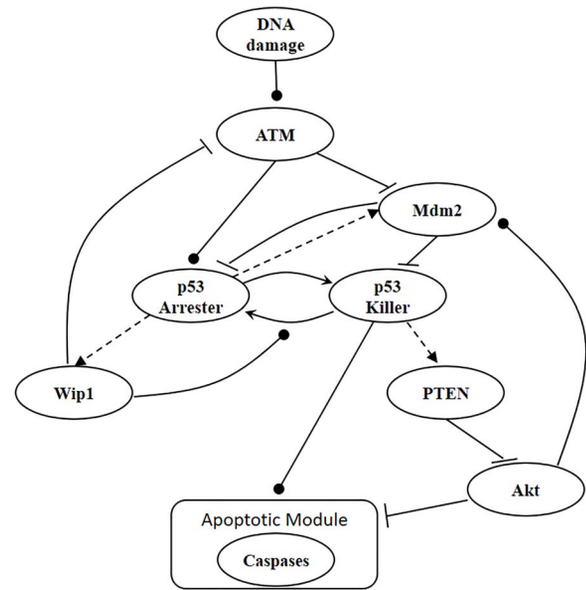


Figure 1. A p53 system model [9]. Arrow-headed dashed lines represent positive transcriptional regulations, arrow-headed solid lines stand for protein transformations, circle-headed solid lines are activatory regulations, and hammer-headed solid lines represent inhibitory regulations. All the molecules and the interactions between them are described in the main body of the paper.

In response to an IR dose, two types of incorrect decisions can be made. One is deciding that the input IR level is high, whereas in fact it is low (deciding H_1 when H_0 is true), which may falsely trigger apoptosis. The other one is deciding that the input IR level is low, whereas in fact it is high (deciding H_0 when H_1 is true), which may result in missing apoptosis. These two erroneous decisions can be called as ‘false alarm’ and ‘miss event’, respectively, and their probabilities can be defined as:

$$\begin{aligned} P_{FA} &= P(\text{deciding } H_1 | H_0), \\ P_M &= P(\text{deciding } H_0 | H_1). \end{aligned} \quad (2)$$

The overall error probability P_E of making decisions is a combination of P_{FA} and P_M :

$$P_E = P(H_0) P_{FA} + P(H_1) P_M, \quad (3)$$

where $P(H_0)$ and $P(H_1)$ are prior probabilities of H_0 and H_1 , respectively. Note that as mentioned in the Introduction section, IR causes DNA damage. Therefore, one can instead formulate the p53-based decision-making process as a binary hypothesis testing on DNA damage being low or high and define the associated false alarm and miss event probabilities accordingly.

The optimal decision-making system which minimizes the above P_E is the one that compares probabilities of observed data under the hypotheses H_0 and H_1 [30]. More precisely, suppose that x is the observation and $p(x|H_0)$ and $p(x|H_1)$ are the conditional probability density functions (PDFs) of x under H_0 and H_1 , respectively. Also consider equi-probable hypotheses, i.e. $P(H_0) = P(H_1) = 0.5$, which is a reasonable assumption in the absence of prior information on the possibilities of H_0 and H_1 . Then, the optimal system decides H_1 if $p(x|H_1) > p(x|H_0)$; otherwise, it decides the H_0 . This means that the hypothesis

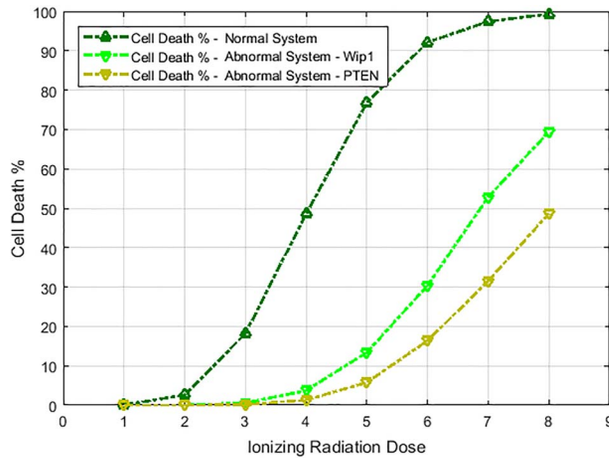


Figure 2. Cell death percentage versus IR dose in both normal and abnormal p53 systems. The dark green curve at the top represents a normal p53 system with no perturbation, whereas the other two curves correspond to p53 systems behaving abnormally due to Wip1 or PTEN perturbations.

with the highest likelihood is decided. This decision is called the maximum likelihood decision [30].

SINGLE CELL DATA OF THE P53 SYSTEM EXPOSED TO IR

To calculate the error probabilities in Equation (2), we use PTEN level as the decision variable because when unreparable DNA damage occurs, the activated p53 triggers pro-apoptotic phosphatase PTEN [27], and PTEN initiates apoptosis [28]. It has also been shown by Hat *et al.* [9] that PTEN is a decent predictor of cell fate. After specifying the decision variable, we use the stochastic simulator of Hat *et al.* [9] to generate 5000 cells for each IR dose. The stochastic simulation has three phases. The first phase is the ‘equilibrium phase’ where we simulate 2 weeks of cell behavior when no IR dose is applied. The second phase is called ‘irradiation phase’ in which 10 minutes of IR dose is applied. The last phase is called ‘relaxation phase’ in which we simulate 72 hours of cell behavior after it is exposed to 10 minutes of IR. When IR dose increases, apoptotic cell percentage increases as well [9] (Fig. 2). For more details on the simulation phases, see supporting files of Hat *et al.* [9]. In order to decide whether a cell is apoptotic or not, we check active caspase level 72 hours after the irradiation phase and compare it with the threshold of 0.5×10^5 suggested in Hat *et al.* [9]. Cells with the level of active caspase higher (or lower) than the threshold of 0.5×10^5 are considered to be apoptotic (or surviving).

The data of normal cells includes eight sets of PTEN levels in 5000 cells, which correspond to eight doses of IR = 1, 2, 3, 4, 5, 6, 7 and 8 Gy. Here Gy stands for Gray, the unit of radiation dose, and 1 Gy is 1 Joule of energy absorbed by 1 kg of tissue. We focus our analysis on low IR versus high IR hypothesis testing, to see how accurately it can be decided whether the applied radiation level is low or high. We consider IR = 1 Gy as the low dose, whereas the higher dose can be IR = 2, 3, 4, 5, 6, 7 or 8 Gy. More specifically, scenarios in which signaling outcomes are analyzed are 1 vs. 2 Gy, 1 vs. 3 Gy, 1 vs. 4 Gy, 1 vs. 5 Gy, 1 vs. 6 Gy, 1 vs. 7 Gy and 1 vs. 8 Gy. We quantitatively study in which of these scenarios more erroneous decisions are made. We also determine to what extent decision between responses to low and high IR levels depends on the input IR separation. We conduct these studies by computing

the optimal decision threshold in each scenario using the PTEN data, following the maximum likelihood principle that provides the best decisions, i.e. smallest decision error probabilities. We also compute numerical values of the decision error probabilities using the PTEN data.

In addition to the analysis of erroneous decision-making and incorrect signaling outcomes in normal cells mentioned above, we analyze them in abnormal cells as well, where there is a dysfunctional molecule in the p53 system. Wip1 is one of the key regulatory pro-survival phosphatases [23] in the p53 system (Fig. 1). If the DNA damage can be repaired, then Wip1 expression returns the cell to the pre-stress state from cell-cycle arrest [23, 32]. It has been observed that elevated Wip1 level exists in multiple human cancer types such as breast, lung, pancreas, bladder and liver cancer [33–38]. Therefore, to obtain abnormal cells, we generate cells with increased Wip1 synthesis rate. In normal cells, Wip1 synthesis rate is about 0.1 [9], and here we increase it to 0.15, a 50% increase, to reproduce abnormality. This increase in the Wip1 synthesis rate causes a significant decrease in the cell death percentage (Fig. 2), which can be considered as an abnormal cell state. In addition to Wip1, we analyze abnormal cellular state caused by PTEN abnormalities. It has been observed that attenuated PTEN levels exist in MCF-7, a noninvasive form of human breast cancer cells [39]. Therefore, it is of interest to see how the abnormal PTEN level affects signaling outcomes in the p53 system. To study this, we generate abnormal cells by decreasing PTEN synthesis rate. In healthy cells, the PTEN synthesis rate is about 0.03 [9]. Here we decrease it to 0.015, a 50% decrease, to reproduce abnormality. We observe a considerable decrease in the cell death percentage (Fig. 2), representing an abnormal cellular state.

UNIVARIATE ANALYSIS: METHODS FOR COMPUTING DECISION THRESHOLDS AND DECISION ERROR RATES USING ‘SINGLE’ TIME POINT MEASUREMENTS IN INDIVIDUAL CELLS

In this section, we analyze PTEN levels of 5000 cells measured in 72 hours after the irradiation phase. It has been observed that PTEN levels of both apoptotic and surviving cells become very distinct in 72 hours after 10 minutes of IR application [9] (decision analysis based on PTEN levels at other time instants and multiple time instants are presented in other sections).

Histograms of natural logarithm, \ln , of PTEN levels for IR = 1 and 2 Gy data sets and IR = 1 and 8 Gy data sets are shown in Fig. 3A and C, respectively. As presented in Fig. 3B and D, Gaussian PDFs whose means and variances are estimated from the data reasonably represent the histograms. This indicates that the PTEN data can be reasonably approximated by lognormal PDF. Due to the mathematical convenience of working with Gaussian PDFs and variables, especially for multivariate analysis of multiple time point data discussed later, we continue working with the logarithm of the PTEN data. Let $x = \ln(\text{PTEN})$ be the Gaussian variable of interest with mean μ and variance σ^2 , i.e., $x \sim \mathcal{N}(\mu, \sigma^2)$ where \mathcal{N} stands for the following normal or Gaussian PDF:

$$p(x) = \left(2\pi\sigma^2\right)^{-1/2} \exp\left[-(x - \mu)^2 / \left(2\sigma^2\right)\right].$$

The Gaussian PDFs shown in Fig. 3 are indeed the conditional PDFs $p(x|H_0)$ and $p(x|H_1)$ under the hypotheses H_0 and H_1

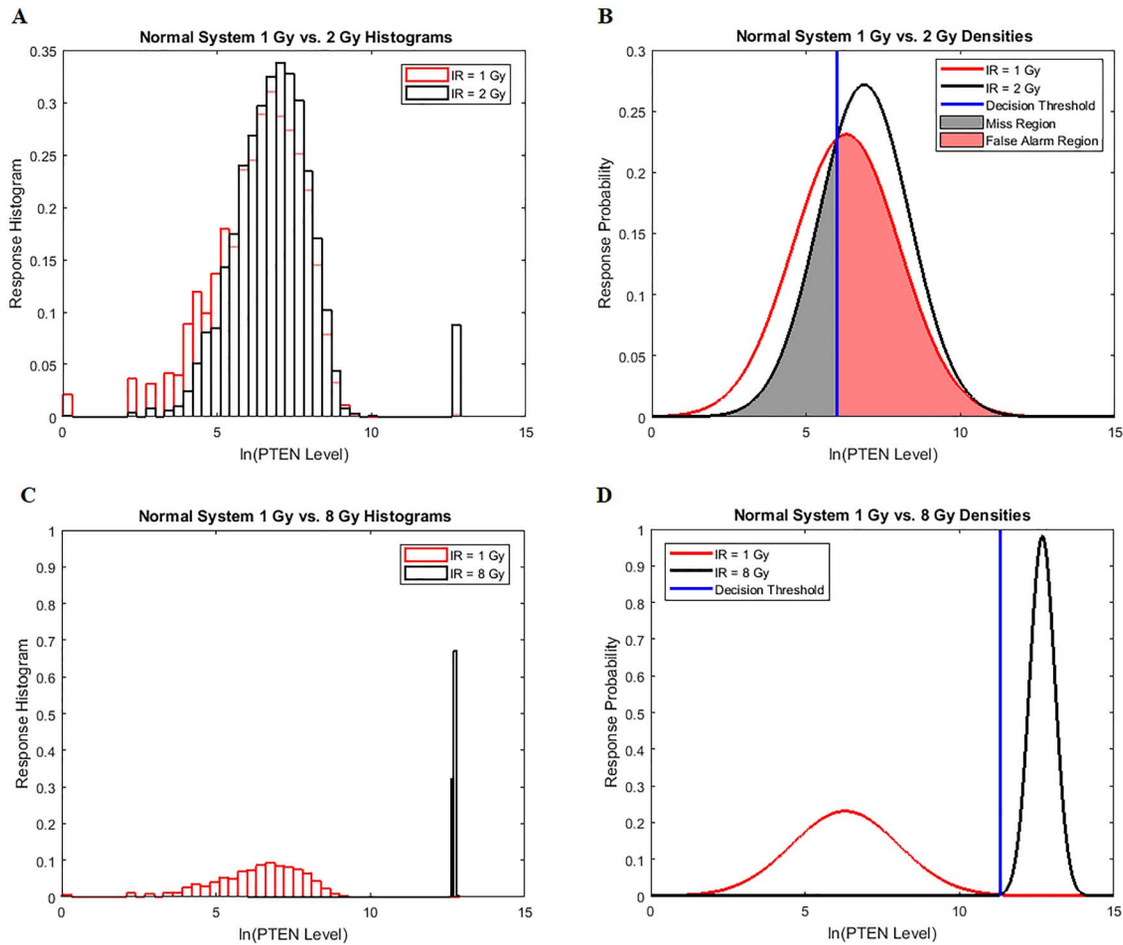


Figure 3. Univariate decision-making and signaling outcome analysis in the normal p53 system based on PTEN response distributions. (A) Histograms of PTEN levels of cells under IR = 1 and 2 Gy doses. (B) Gaussian PDFs for PTEN levels of cells under IR = 1 and 2 Gy doses, together with the optimal maximum likelihood decision threshold which minimizes the total decision error probability. (C) Histograms of PTEN levels of cells under IR = 1 and 8 Gy doses. (D) Gaussian PDFs for PTEN levels of cells under IR = 1 and 8 Gy doses, together with the optimal maximum likelihood decision threshold which minimizes the total decision error probability.

defined earlier in Equation (1). For example, in Fig. 3B, H_0 and H_1 correspond to IR = 1 and 2 Gy doses, respectively, and the red and black curves in there are the conditional PDFs $p(x|H_0)$ and $p(x|H_1)$, respectively.

The optimal maximum likelihood decision-making system

Recall our two hypotheses previously defined in (1). The optimal decision-maker, which minimizes the overall error probability P_E in (3), compares the conditional likelihood ratio $L(x) = p(x|H_1)/p(x|H_0)$ with the ratio $\gamma = P(H_0)/P(H_1)$ [1]. The system decides H_1 if $L(x) > \gamma$. If the hypotheses are equi-probable, i.e. $P(H_0) = P(H_1) = 0.5$, then the optimal system decides H_1 if $p(x|H_1) > p(x|H_0)$.

The optimal decision threshold

To find the optimal decision threshold, we need to solve the equation $L(x) = \gamma$, i.e. $P(H_1)p(x|H_1) = P(H_0)p(x|H_0)$, for x . When H_0 and H_1 are equi-probable, the threshold equation to be solved simplifies to $L(x) = 1$, i.e. $p(x|H_1) = p(x|H_0)$.

The decision error probabilities

Once the optimal decision threshold is determined, it can be used to compute false alarm and miss decision error probabilities, by integrating the conditional PDFs of data over error regions. More specifically, using the conditional PDFs $p(x|H_0)$ and $p(x|H_1)$ representing the response probabilities of the ln of PTEN levels under the two hypotheses, Equation (2) can be written as [1]:

$$P_{FA} = \int_{x \in \text{false alarm region}} p(x|H_0) dx, \quad (4)$$

$$P_M = \int_{x \in \text{miss region}} p(x|H_1) dx. \quad (5)$$

The false alarm region in (4) is defined by $\{x : p(x|H_1) > p(x|H_0)\}$ when H_0 is true, whereas the miss region in (5) is defined by $\{x : p(x|H_0) > p(x|H_1)\}$ when H_1 is true. By substituting P_{FA} and P_M in Equation (3), the overall error probability P_E can be obtained.

The Gaussian data model to compute the optimal decision threshold

Here we focus on Fig. 3B as an example, where two Gaussian PDFs are shown for $x = \ln(\text{PTEN})$, the natural logarithm of PTEN levels in the two data sets of IR = 1 and 2 Gy, with each data set consisting of 5000 cells. Let $\mathcal{N}(\mu_0, \sigma_0^2)$ and $\mathcal{N}(\mu_1, \sigma_1^2)$ represent the Gaussian PDFs that correspond to the IR = 1 and 2 Gy data sets, respectively, where (μ_0, σ_0^2) and (μ_1, σ_1^2) are mean/variance pairs estimated from their associated data sets. The optimal maximum likelihood decision threshold in Fig. 3B is at the intersection of the two PDFs and can be computed by solving the equation $p(x|H_0) = p(x|H_1)$ written below:

$$(2\pi\sigma_0^2)^{-1/2} \exp\left[-(x - \mu_0)^2 / (2\sigma_0^2)\right] = (2\pi\sigma_1^2)^{-1/2} \exp\left[-(x - \mu_1)^2 / (2\sigma_1^2)\right]. \quad (6)$$

By multiplying both sides by $(2\pi\sigma_0^2)^{1/2} \exp[(x - \mu_1)^2 / (2\sigma_1^2)]$ and then taking natural logarithm of both sides, (6) can be written in the following quadratic equation form [1]:

$$(\sigma_0^2 - \sigma_1^2)x^2 + 2(\sigma_1^2\mu_0 - \sigma_0^2\mu_1)x + \sigma_0^2\mu_1^2 - \sigma_1^2\mu_0^2 - 2\sigma_0^2\sigma_1^2 \ln(\sigma_0/\sigma_1) = 0. \quad (7)$$

Equation (7) is derived assuming our hypotheses are equiprobable, i.e. $P(H_0) = P(H_1) = 0.5$, as mentioned before. The solution of Equation (7) gives the optimal decision threshold PTEN_{th} , located at the intersection of the two PDFs for IR = 1 and 2 Gy doses in Fig. 3B (the italic style is adopted to clarify that the threshold is related to the logarithm of PTEN data). Interestingly, for equal variances, the solution of Equation (7) for the optimal decision threshold simplifies to the average of the means, i.e. $(\mu_0 + \mu_1)/2$, which intuitively makes sense. For other prior probabilities and PDF models, the optimal threshold can be obtained similarly, by solving the equation $P(H_0)p(x|H_0) = P(H_1)p(x|H_1)$ for x .

The Gaussian data model to compute the decision error probabilities

Using the PTEN_{th} obtained by solving Equation (7) and using the Gaussian PDFs, Equations (4) and (5) for the false alarm and miss error probabilities can be written as:

$$P_{\text{FA}} = \int_{\text{PTEN}_{\text{th}}}^{\infty} p(x|H_0) dx = Q\left(\frac{\text{PTEN}_{\text{th}} - \mu_0}{\sigma_0}\right), \quad (8)$$

$$P_{\text{M}} = \int_{-\infty}^{\text{PTEN}_{\text{th}}} p(x|H_1) dx = Q\left(\frac{\mu_1 - \text{PTEN}_{\text{th}}}{\sigma_1}\right), \quad (9)$$

where $Q(\eta)$ is tail probability of the standard Gaussian PDF :

$$Q(\eta) = (2\pi)^{-1/2} \int_{\eta}^{\infty} \exp(-u^2/2) du.$$

Equation (8) represents area of the pink region in Fig. 3B under the tail of the IR = 1 Gy PDF, beyond the PTEN_{th} threshold. In this region of $x > \text{PTEN}_{\text{th}}$, we have $p(x|H_1) > p(x|H_0)$, while H_0 is true. This is the ‘false alarm’ region for which we have

computed $P_{\text{FA}} = 0.57$ in Fig. 3B. On the other hand, Equation (9) represents the area of the gray region in Fig. 3B under the tail of the IR = 2 Gy PDF, below the PTEN_{th} threshold. In this region of $x < \text{PTEN}_{\text{th}}$, we have $p(x|H_0) > p(x|H_1)$, while H_1 is true. This is the ‘miss’ region for which we have computed $P_{\text{M}} = 0.28$ in Fig. 3B. After computing P_{FA} and P_{M} , we can now compute the overall error probability P_{E} using Equation (3), which results in $P_{\text{E}} = (P_{\text{FA}} + P_{\text{M}})/2 = 0.43$. Similarly, by computing Equations (8) and (9) for the 1 vs. 8 Gy scenario, we obtain $P_{\text{E}} = 0.001$ (Fig. 3D). Based on the results of 1 vs. 2 Gy and 1 vs. 8 Gy decision scenarios, it can be concluded that when the difference between the two applied IR doses increases, the overall decision error probability P_{E} decreases. This is mainly because the two response PDFs become more distinct with less overlap, as the difference between the two applied IR doses increases.

For some cases such as 1 vs. 3, 4, 5 and 6 Gy IR doses, some data sets need to be modeled by a mixture of Gaussian PDFs due to the bistable behavior of p53 system and hence cells’ bimodal histograms. Still the same underlying theory and proposed framework hold. Nevertheless, in what follows we explain how to determine the optimal decision thresholds and how to compute the decision error probabilities when using a mixture model, for the 1 vs. 4 Gy scenario.

Histograms of natural logarithm of PTEN levels for IR = 1 and 4 Gy data sets are shown in Fig. 4A. We notice that while 1 Gy data histogram is unimodal, histogram of 4 Gy data is bimodal. Therefore, for the 1 Gy data, we use a single Gaussian PDF as before, whereas for the 4 Gy data, we utilize a mixture of two Gaussian PDFs. More specifically, we consider $\mathcal{N}(\mu_0, \sigma_0^2)$ for H_0 to represent the single Gaussian PDF that corresponds to the IR = 1 Gy data, whereas we use $\xi \mathcal{N}(\mu_{11}, \sigma_{11}^2) + (1 - \xi) \mathcal{N}(\mu_{12}, \sigma_{12}^2)$ for H_1 , with $0 \leq \xi \leq 1$ being the mixing parameter, to represent the mixture of two Gaussian PDFs which correspond to the IR = 4 Gy data set. The mean and variance (μ_0, σ_0^2) are estimated from the 1 Gy data, and the associated single Gaussian PDF is shown in Fig. 4B. Furthermore, the means and variances $(\mu_{11}, \sigma_{11}^2)$ and $(\mu_{12}, \sigma_{12}^2)$ and the mixing parameter ξ are estimated from the 4 Gy data using the MATLAB command ‘fitgmdist’ which implements the iterative expectation-maximization algorithm. The resulting mixture of two Gaussian PDFs is shown in Fig. 4B.

Similar to the previous scenarios, the optimal maximum likelihood decision thresholds shown in Fig. 4B for equiprobable hypotheses are at the intersections of the conditional PDFs $p(x|H_0)$ and $p(x|H_1)$, the latter being a Gaussian mixture for the 4 Gy data. Note that here solving the equation $p(x|H_0) = p(x|H_1)$ results in four solutions for x ; that is why there are four decision thresholds, $\text{PTEN}_{\text{th}, i}$, $i = 1, 2, 3, 4$ in Fig. 4B (note that each decision threshold $\text{PTEN}_{\text{th}, i}$ is listed as ‘Decision Threshold i ’ in Fig. 4).

To compute the decision error probabilities, the false alarm and miss probabilities P_{FA} and P_{M} need to be calculated using Equations (4) and (5), respectively. Since there are four decision thresholds in this case, integration has to be performed over multiple regions, which results in lengthy expressions. However, note that as can be seen in Fig. 4B and its zoomed-in view in Fig. 4C, PDFs for low dose (red) and the lower Gaussian mode for the high dose (black) assume very small values as they reach the third threshold. Therefore, their contributions to possible error events around the third and fourth thresholds are negligible (later this is shown numerically). Similarly, given the very small variance of the higher Gaussian mode of the PDF for the high dose, this PDF is substantially different from zero only between the third and fourth thresholds. Consequently, the contribution of the PDF of this mode to possible errors around the third and fourth thresholds is negligible as well. Overall, as just explained,

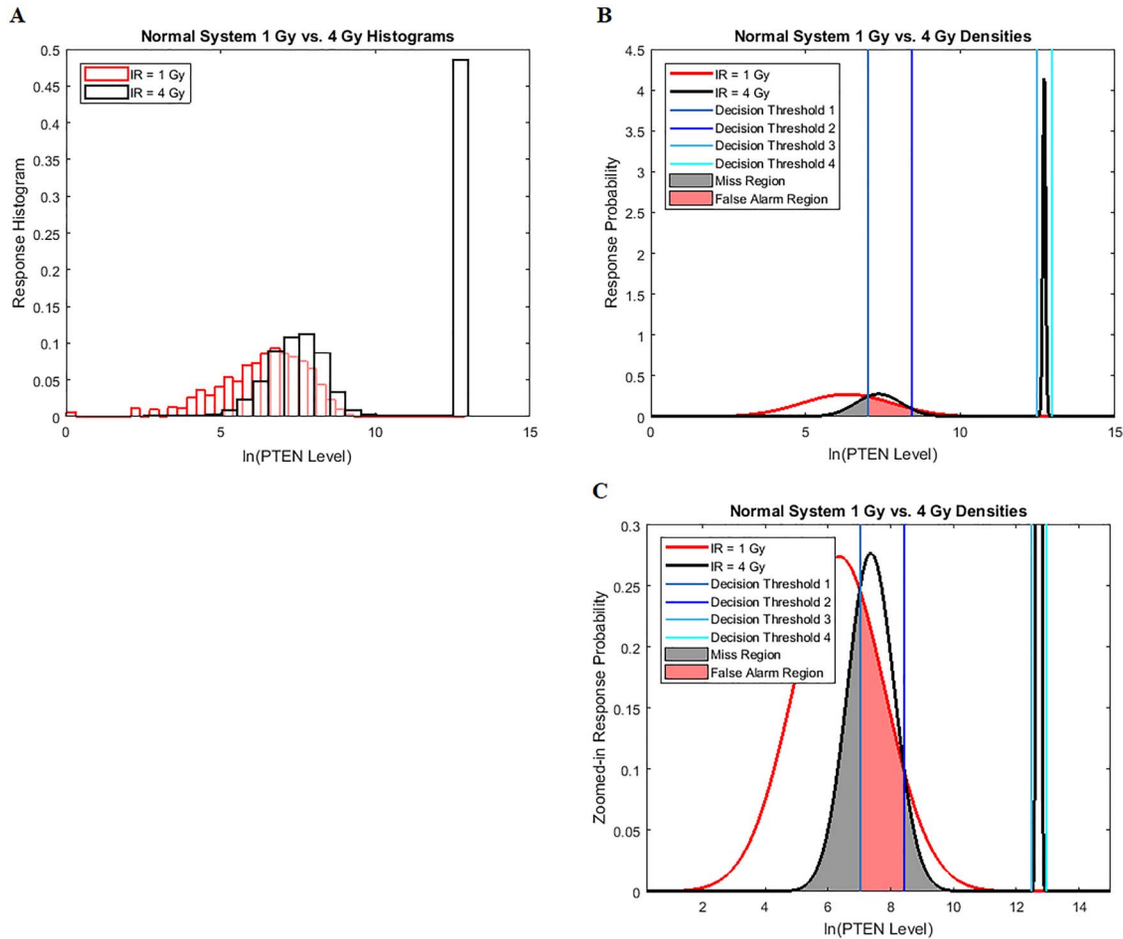


Figure 4. Univariate decision making and signaling outcome analysis in the normal p53 system when a PTEN response distribution is bimodal. (A) Histograms of PTEN levels of cells under IR = 1 and 4 Gy doses. (B) A Gaussian PDF for PTEN levels of cells under IR = 1 Gy and a mixture of two Gaussian PDFs for PTEN levels of cells under IR = 4 Gy doses, together with the optimal maximum likelihood decision thresholds which minimize the total decision error probability. (C) Zoomed-in view of panel B.

optimal decision when $PTEN_{th3} < x < PTEN_{th4}$ is H_1 with no decision error, whereas for $x < PTEN_{th1}$, $PTEN_{th1} < x < PTEN_{th2}$ and $PTEN_{th2} < x < PTEN_{th3}$, optimal decisions are H_0 , H_1 and H_0 , respectively, with the following decision error probabilities:

$$P_{FA} = Q\left(\frac{PTEN_{th1} - \mu_0}{\sigma_0}\right) - Q\left(\frac{PTEN_{th2} - \mu_0}{\sigma_0}\right),$$

$$P_M = \xi \left[Q\left(\frac{\mu_{11} - PTEN_{th1}}{\sigma_{11}}\right) + Q\left(\frac{PTEN_{th2} - \mu_{11}}{\sigma_{11}}\right) \right].$$

The P_{FA} expression corresponds to the pink region in Fig. 4C, whereas the two Q functions in the P_M expression correspond to the two gray regions in Fig. 4C, respectively. Using the data, computed numerical values are $\xi = 0.51$, $P_{FA} = 0.28 - 0.06 \approx 0.22$, $P_M = 0.51 [0.41 + 0.07] \approx 0.25$ and $P_E \approx 0.24$, the last one being calculated using Equation (3).

As an example of a negligible decision error probability around the third and fourth thresholds mentioned earlier, consider the area under the red Gaussian PDF $p(x|H_0)$ in Fig. 4C for $PTEN_{th3} < x < PTEN_{th4}$. While not visible due to being very small, it can be understood that the aforementioned area is a false alarm probability of deciding H_1 , although H_0 is true. The numerical value of this false alarm probability is: $Q((PTEN_{th3} - \mu_0)/\sigma_0) - Q((PTEN_{th4} - \mu_0)/\sigma_0) = 1.2 \times 10^{-5} - 2.8 \times 10^{-6} \approx 0$, which

is negligible compared to $P_{FA} \approx 0.22$ calculated in the previous paragraph.

Abnormal p53 systems

To see how an abnormality in the p53 system affects the decision-making and signaling outcomes, we calculate P_E values when Wip1 synthesis rate is elevated by 50% from 0.1 to 0.15 (Fig. 5), as mentioned previously. As suggested by Habibi et al. [1], decision thresholds are modeled to be those of the normal cells. This implies that abnormal cells are not aware of the abnormality and therefore erroneously use the previous threshold. As we see later, this increases decision error probabilities, a behavior that can be anticipated from abnormal cells. Using Equations (3), (8) and (9), P_{FA} , P_M and P_E are computed: $P_E = 0.44$ is obtained for 1 vs. 2 Gy scenario (Fig. 5A), and $P_E = 0.16$ is obtained for 1 vs. 8 Gy scenario (Fig. 5B). Compared to the normal system results, the overall error probability is significantly higher for the abnormal system under the 1 vs. 8 Gy scenario (we observe that $P_E = 0.001$ of normal cells markedly increases to $P_E = 0.16$ in abnormal cells). The reason is that when the Wip1 synthesis rate is increased, the two response PDF curves significantly overlap (notice the overlap between the left-side component of the IR = 8 Gy PDF with the IR = 1 Gy

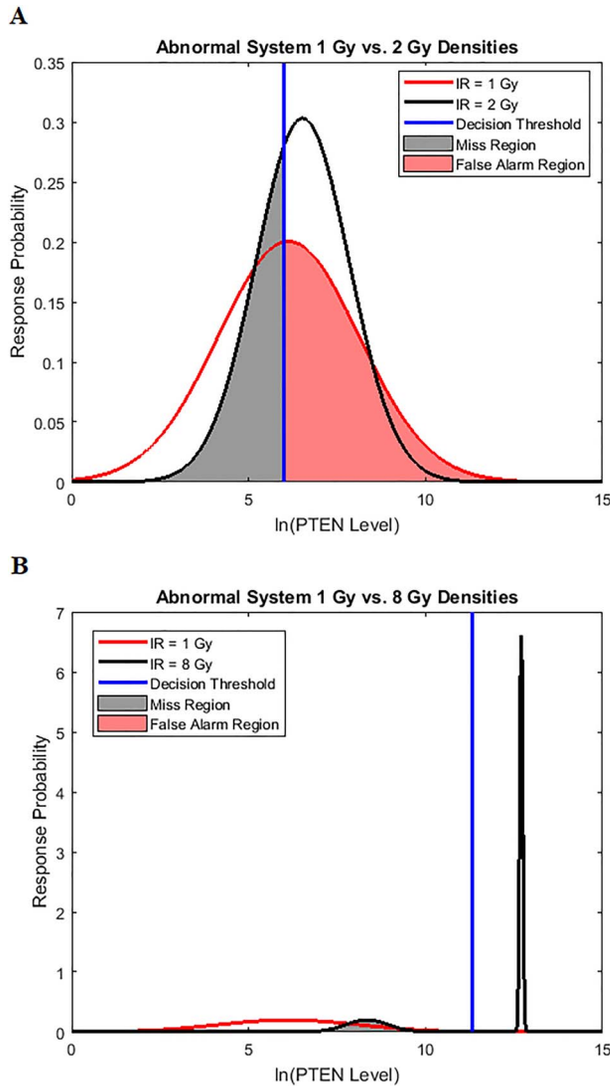


Figure 5. Univariate decision-making and signaling outcome analysis in an abnormal p53 system, with increased Wip1 synthesis rate, based on PTEN response distributions. (A) Gaussian PDFs for PTEN levels of abnormal cells under IR = 1 and 2 Gy doses, together with the decision threshold of normal cells. This implies that in abnormal cells the previous decision threshold is erroneously used [1]. As discussed later, this increases decision error probabilities, a behavior that can be anticipated from abnormal cells. (B) A Gaussian PDF for PTEN levels of abnormal cells under IR = 1 Gy dose and a mixture of two Gaussian PDFs for PTEN levels of abnormal cells under IR = 8 Gy dose, together with the decision threshold of normal cells.

PDF in Fig. 5B). This is while in normal cells they had almost no overlap (Fig. 3D).

Similarly, we compute error probabilities for the other abnormal p53 system we mentioned previously, generated by the PTEN synthesis rate reduced from 0.03 to 0.015 (50% reduction). Error probabilities for this abnormality for all different radiation exposure scenarios of 1 vs. 2 Gy up to 1 vs. 8 Gy are shown in Fig. 6. For comparison, error probabilities for the Wip1-perturbed abnormal p53 system and also the normal p53 system are provided in Fig. 6 as well. We observe that as the difference between the two applied IR doses increases, decision error probability in normal cells drops significantly. This is while, in abnormal cells, decision error probabilities remain high. These signaling outcomes might be correlated with the observation that cell death percentages

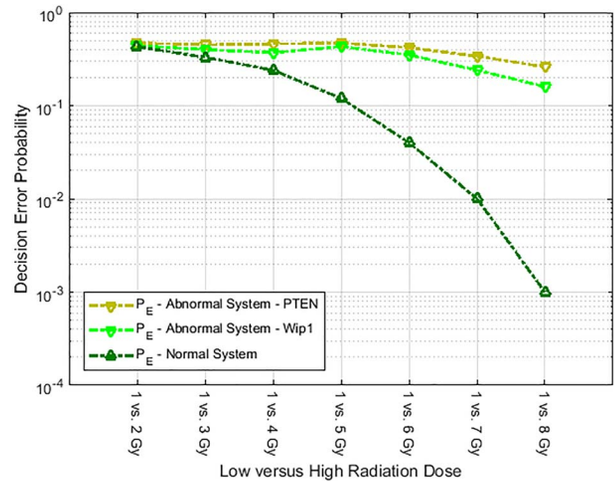


Figure 6. Decision error probabilities for several low IR versus high IR scenarios. The ‘Abnormal System—PTEN’ legend refers to a p53 system whose PTEN synthesis rate is decreased by 50%, compared to its nominal value. The ‘Abnormal System—Wip1’ legend refers to a p53 system whose Wip1 synthesis rate is increased by 50%, compared to its nominal value. Smaller decision error probabilities in the normal system are noteworthy.

in abnormal systems are considerably lower than the normal system, even when the radiation dose increases (Fig. 2). This could indicate that abnormal cells do not respond to IR levels properly and, hence, decisions and signaling outcomes affecting apoptosis and survival become more erroneous. Care should be taken that these specific observations are based on the low versus high IR, e.g. d_0 vs. d_1 IR hypothesis testing formulation, where the low IR dose is fixed to 1 Gy ($d_0 = 1$ Gy) and the high IR dose is ranging from 2 up to 8 Gy ($d_1 = 2, 3, \dots, 8$ Gy) in the p53 system, that is considered in this paper as an example. These observations may not be generalized to other selections of the low d_0 and high d_1 IR doses or other hypothesis testing formulations, case studies or signaling networks. However, the proposed framework and its analytical tools, whose introduction has been the main goal of this paper, can be similarly used.

DECISION AND SIGNALING OUTCOME ANALYSIS USING ROC CURVES

In this subsection, we show how to analyze performance of a decision-maker using ROC curves. The ROC curve is developed to visualize the performance of decision-making systems [30, 40] and is a graph of probability of detection, $P_D = 1 - P_M$, versus the probability of false alarm, P_{FA} . In Fig. 7 we present ROC curves for both the normal p53 system (Fig. 7A) and the abnormal p53 system (Fig. 7B) whose Wip1 synthesis rate is elevated, for these two low versus high IR decision-making scenarios: 1 vs. 2 Gy and 1 vs. 8 Gy. The theoretical ROC curves in Fig. 7 are graphed using the false alarm and miss decision error probability formulas in Equations (8) and (9), respectively, with μ, σ and the thresholds estimated from the data. The empirical ROC curves in Fig. 7 are graphed by using the data sets directly, using the MATLAB command ‘percurve’. We observe that the theoretical and empirical ROCs are nearly the same. Therefore, in what follows, we focus on the theoretical ROC curves, to explain concepts and results.

A ROC curve is above a 45° diagonal line [30], the gray dashed line in Fig. 7. In our study it represents the worst possible decision-maker, i.e. a decision-making system that does not

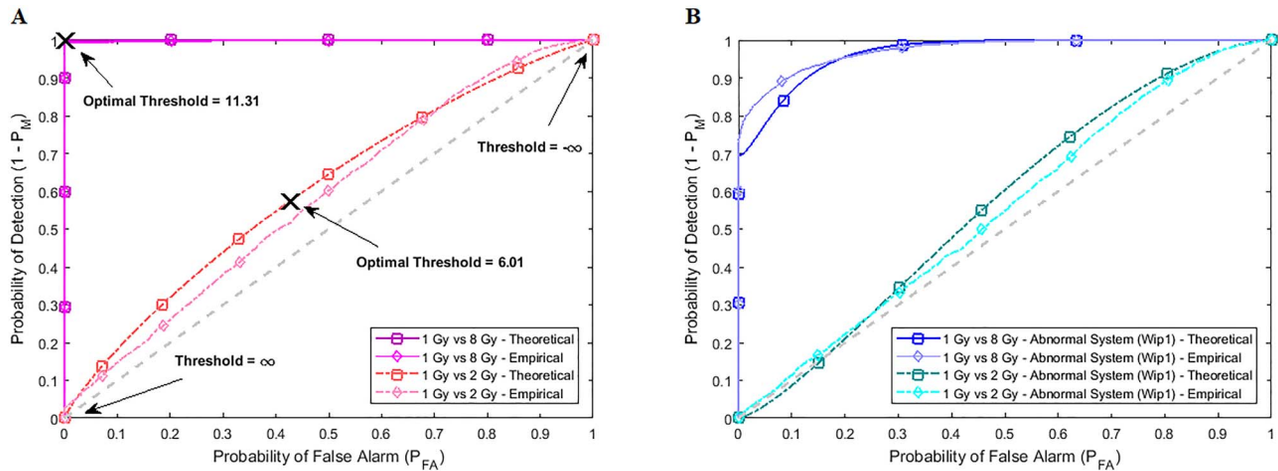


Figure 7. Empirical and theoretical ROC curves for both normal and abnormal p53 systems. The theoretical ROC curves labeled by \square are obtained from the Gaussian and mixture of Gaussian data models and formulas whose parameters are estimated from the data, whereas the empirical ROC curves labeled by \diamond are obtained directly from the data. We observe that the theoretical and empirical ROCs are nearly the same. Note that Threshold = $\ln(\text{PTEN Level})$ in the figures. (A) ROC curves of 1 vs. 2 Gy and 1 vs. 8 Gy radiation scenarios for the normal system. (B) ROC curves of 1 vs. 2 Gy and 1 vs. 8 Gy radiation scenarios for the Wip1-perturbed abnormal system.

use the data and instead randomly decides if the applied IR dose is low or high, by just flipping a coin. The 45° line is indeed a reference to judge the performance of a decision-making system. A ROC curve far away from the 45° reference line indicates a good decision-maker. Each point on a ROC curve represents a (P_{FA}, P_D) pair that corresponds to a certain decision threshold. Other properties of ROC curves can be found in Van Trees *et al.* [40]. The ‘x’ marks in Fig. 7A show the optimal (P_{FA}, P_D) points that correspond to the optimal decision thresholds shown in Fig. 3B and D, previously computed using Equation (7) for the 1 vs. 2 Gy and 1 vs. 8 Gy scenarios, respectively.

Based on the normal p53 system ROC curves in Fig. 7A, we observe that decisions are made better under the 1 vs. 8 Gy scenario, because of its ROC curve being very far from the 45° reference line, compared to the 1 vs. 2 Gy case whose ROC curve is much closer to the 45° reference line. This finding supports our results presented in Fig. 6, showing the smaller decision error probability of 0.001 for 1 vs. 8 Gy, compared to the larger decision error probability of 0.43 for 1 vs. 2 Gy. ROC curves also show that abnormalities in the p53 system can cause decision precision loss. Comparing the normal (Fig. 7A) and abnormal system ROC curves (Fig. 7B), we observe that the abnormal system ROC curves are closer to the 45° reference line, meaning that more erroneous decisions are made, when there is an abnormality in the system.

BIVARIATE ANALYSIS: METHODS FOR COMPUTING DECISION THRESHOLDS AND DECISION ERROR RATES USING ‘TWO’ TIME POINT MEASUREMENTS IN INDIVIDUAL CELLS

In this section, we analyze PTEN levels of 5000 cells measured in 1 hour and 30 hours after the irradiation phase. Using two variables instead of one allows to study the effect of temporal dynamical changes on decision-making and signaling outcomes and paves the way for analyzing decisions based on multiple time point data. Suppose x and y represent the $\ln(\text{PTEN})$ levels in 1 hour and 30 hours, respectively, after radiation. Joint Gaussian

PDF for x and y can be written as [41]:

$$p(x, y) = \frac{1}{2\pi\sigma_x\sigma_y\sqrt{1-\rho^2}} \exp\left(-\frac{1}{2(1-\rho^2)} \left[\frac{(x-\mu_x)^2}{\sigma_x^2} - \frac{2\rho(x-\mu_x)(y-\mu_y)}{\sigma_x\sigma_y} + \frac{(y-\mu_y)^2}{\sigma_y^2} \right] \right), \quad (10)$$

where (μ_x, σ_x^2) and (μ_y, σ_y^2) are means and variances of x and y and ρ is the correlation coefficient between x and y . Bivariate conditional likelihood ratio is given by $L(x, y) = p(x, y|H_1)/p(x, y|H_0)$, and the optimal decision-maker which minimizes the overall error probability P_E compares $L(x, y)$ with the ratio $\gamma = P(H_0)/P(H_1)$. The system decides the H_1 if $L(x, y) > \gamma$. If the hypotheses are equi-probable, i.e. $P(H_0) = P(H_1) = 0.5$, then the optimal system decides the H_1 if $p(x, y|H_1) > p(x, y|H_0)$. To find the optimal decision threshold curve, we need to solve the equation $L(x, y) = \gamma$, i.e. $P(H_1)p(x, y|H_1) = P(H_0)p(x, y|H_0)$, for x and y . When H_0 and H_1 are equi-probable, the threshold equation to be solved simplifies to $L(x, y) = 1$, i.e. $p(x, y|H_1) = p(x, y|H_0)$. To find false alarm and miss probabilities, Equations (4) and (5) can be extended to two variables as follows:

$$P_{FA} = \iint_{(x,y) \in \text{false alarm region}} p(x, y|H_0) dx dy, \quad (11)$$

$$P_M = \iint_{(x,y) \in \text{miss region}} p(x, y|H_1) dx dy, \quad (12)$$

where $\{x, y : p(x, y|H_1) > p(x, y|H_0)\}$ defines the false alarm region when H_0 is true and $\{x, y : p(x, y|H_0) > p(x, y|H_1)\}$ specifies the miss region when H_1 is true. After computing P_{FA} and P_M , the overall decision error probability P_E can be calculated using Equation (3).

As an example, here we focus on Fig. 8A, where two bivariate Gaussian PDFs are shown for $x = \ln(\text{PTEN at the 1st hour})$ and $y = \ln(\text{PTEN at the 30th hour})$, logarithms of PTEN levels in the two data sets of IR = 1 and 2 Gy, with each data set consisting of 5000 cells. The mean and variance parameters of each bivariate response PDF are estimated from the associated data set. The overlap between the two bivariate PDFs in response to IR = 1 and

2 Gy can be better seen in the top view shown in Fig. 8B. This figure also demonstrates that the decision threshold between the two PDFs is going to be a curve in the x - y plane, where the two PDFs intersect. The equation for this optimal threshold curve which minimizes the total decision error probability is given by $L(x, y) = 1$, where L is the bivariate conditional likelihood ratio defined previously. This decision threshold curve $curve_{th}$ is shown together with contour plots of the two bivariate PDFs in Fig. 8C. To compute the decision error probabilities using the decision threshold $curve_{th}$, Equations (11) and (12) for the false alarm and miss error probabilities can be written as:

$$P_{FA} = \int_{x=-\infty}^{\infty} \int_{y=curve_{th}}^{\infty} p(x, y|H_0) dy dx, \quad (13)$$

$$P_M = \int_{x=-\infty}^{\infty} \int_{y=-\infty}^{curve_{th}} p(x, y|H_1) dy dx. \quad (14)$$

After computing the integrals in Equations (13) and (14) numerically, we obtain $P_{FA} = 0.24$ and $P_M = 0.26$. Upon their substitution in Equation (3) and with equi-probable hypotheses, we obtain $P_E = 0.25$.

To compare the above two time point decision with individual one time point decisions, we compute decision error probabilities based on the first hour data and the 30th hour data, individually, for the IR = 1 vs. 2 Gy scenario. We obtain $P_E = 0.5$ and $P_E = 0.27$ for individual univariate decisions in 1 hour and 30 hours after the radiation, respectively. We observe that the bivariate decision offers significant improvement over the 1 hour decision and slight improvement over the 30 hour decision. Univariate decision error probabilities at different time points are discussed in the next section, as well as how multivariate decision error probability changes, as the data of more time points are added to the decision process in a sequential manner.

MULTIVARIATE ANALYSIS: METHODS FOR COMPUTING DECISION THRESHOLDS AND DECISION ERROR RATES USING ‘MULTIPLE’ TIME POINT MEASUREMENTS IN INDIVIDUAL CELLS

In this section, we further study the effect of system dynamics on decision-making and signaling outcomes, by considering multiple time point data. More specifically, we consider PTEN levels of 5000 cells measured in 1, 10, 20, 30, 40, 50, 60 and 70 hours after the irradiation phase. Let ω be an $N \times 1$ column vector that represents the $\ln(\text{PTEN})$ levels at a subset or all of the aforementioned time instants. Joint Gaussian PDF for all the decision variables in ω can be written as [31, 40]:

$$p(\omega) = \frac{1}{(2\pi)^{N/2} |\Sigma|^{1/2}} \exp \left[-\frac{1}{2} (\omega - \mu)^T \Sigma^{-1} (\omega - \mu) \right], \quad (15)$$

where μ is the $N \times 1$ mean vector, Σ is the $N \times N$ covariance matrix, $|\Sigma|$ and Σ^{-1} denote the determinant and inverse of Σ , respectively, and T represents matrix transpose. This multivariate Gaussian or normal PDF for the decision vector ω can be symbolically shown by $\omega \sim \mathcal{N}(\mu, \Sigma)$. For $N = 2$, Equation (15)

simplifies to the bivariate PDF in Equation (10), such that

$$\omega = \begin{bmatrix} x \\ y \end{bmatrix}, \mu = \begin{bmatrix} \mu_x \\ \mu_y \end{bmatrix}, \Sigma = \begin{bmatrix} \sigma_x^2 & \rho\sigma_x\sigma_y \\ \rho\sigma_x\sigma_y & \sigma_y^2 \end{bmatrix}.$$

Computation of the decision error probabilities using multiple decision variables can be accomplished using discriminant functions [31, 40]:

$$g_i(\omega) = \ln p(\omega|H_i) + \ln P(H_i), i = 0, 1, \quad (16)$$

where $p(\omega|H_i) \sim \mathcal{N}(\mu_i, \Sigma_i)$ and i is index of the discriminant function associated with the hypothesis H_i . In our case we have $i = 0, 1$, referring to our two hypotheses in Equation (1). For any hypothesis H_i , substitution of (15) in (16) simplifies its discriminant function to

$$g_i(\omega) = -\frac{1}{2} (\omega - \mu_i)^T \Sigma_i^{-1} (\omega - \mu_i) - \frac{N}{2} \ln (2\pi) - \frac{1}{2} \ln |\Sigma_i| + \ln P(H_i), i = 0, 1. \quad (17)$$

Using the discriminant functions in (17) and for a given ω , the optimal decision-making system decides the H_0 if $g_0(\omega) > g_1(\omega)$, and decides the H_1 if $g_1(\omega) > g_0(\omega)$. The false alarm probability P_{FA} is the probability of deciding H_1 , i.e. $g_1(\omega) > g_0(\omega)$, whereas in fact H_0 is true. On the other hand, the miss probability P_M is the probability of deciding the H_0 , i.e. $g_0(\omega) > g_1(\omega)$, although indeed H_1 is true. Computing P_{FA} and P_M using multivariate PDFs directly entail multivariate integrations over regions defined by decision surfaces. Given the complexities of such computations, as a simpler alternative, we calculate P_{FA} and P_M using the data directly, by counting the number of times that false alarm and miss event occur, respectively, after comparing the discriminant function values $g_1(\omega)$ and $g_0(\omega)$ for each ω , and then divide them by the total number of data points. The overall decision error probability P_E can be calculated using Equation (3). Another method for computing P_{FA} and P_M relies on characteristic functions [42].

Single-variable decision-making and signaling outcome analysis as time evolves

To understand how decision-making and signaling outcomes may change over time, first we look at the decision error probability P_E using PTEN levels measured at individual consecutive time instants (Fig. 9A), for the 1 vs. 2 Gy scenario. A noteworthy observation is that the decision error exhibits a minimum value. The minimum occurs in 20 hours after the radiation. This can be visually explained by the amount of overlap of PTEN histograms at each individual time point. For instance, we provide histograms of PTEN levels at the 20th and the 70th hours in Fig. 10, for IR = 1 and 2 Gy doses. We observe that the 20th hour histograms have less overlap than the 70th hour histograms, shown in Fig. 10A and B, respectively, which results in the smaller P_E at the 20th hour in Fig. 9A.

Multivariable decision-making and signaling outcome analysis as time evolves

Now we focus on studying how decision-making works, if data of N time instants are utilized, such that $N = 1, 2, \dots, 8$ (Fig. 9B). In the figure, $N = 1$ means the PTEN data of the first hour, $N = 2$

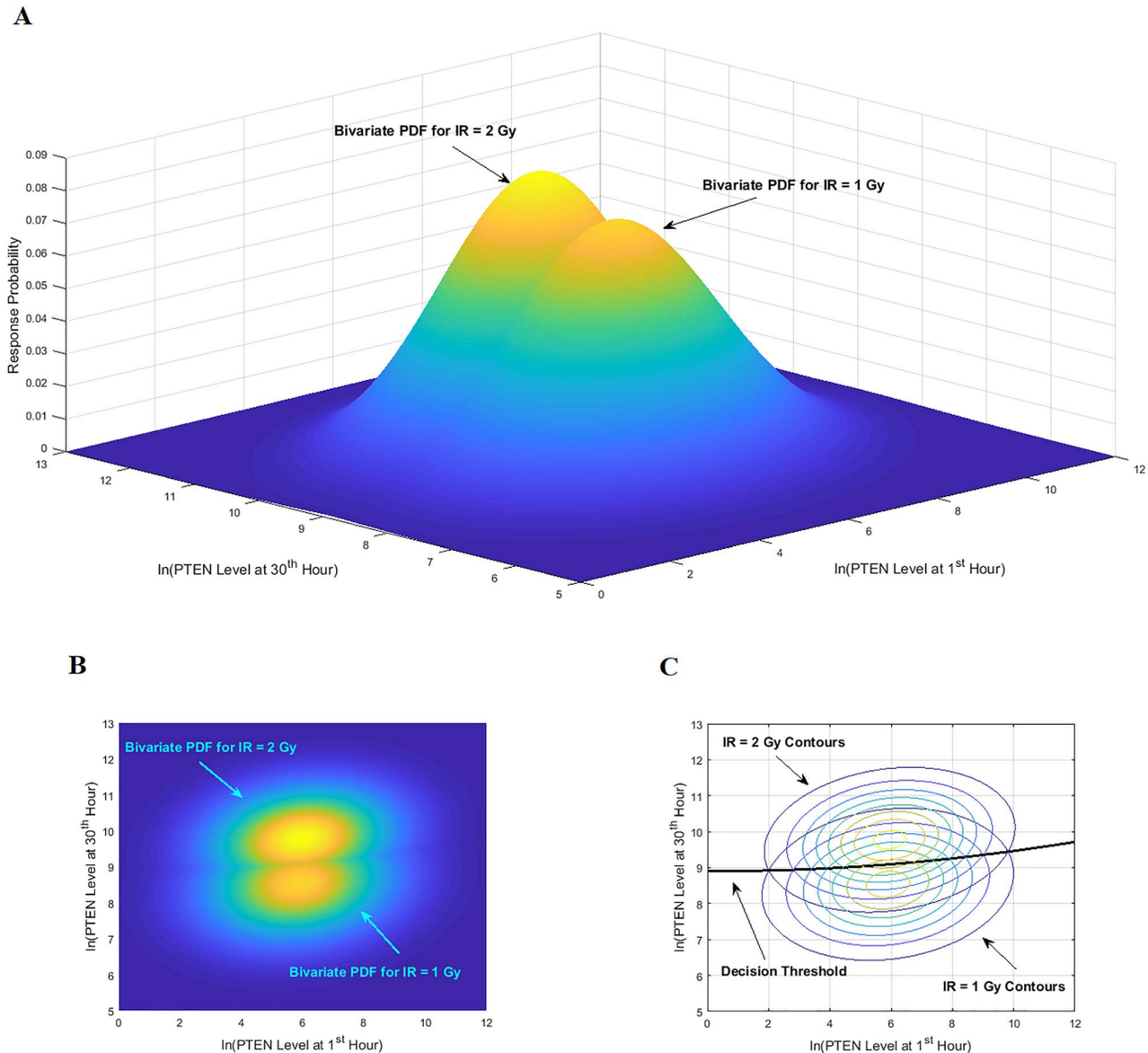


Figure 8. Bivariate decision-making and signaling outcome analysis in the normal p53 system based on PTEN response distributions. (A) Bivariate Gaussian PDFs for PTEN levels of cells at the first hour and the 30th hour, under IR = 1 and 2 Gy doses. (B) Top view of the two bivariate Gaussian PDFs. (C) Top contour view of the two bivariate Gaussian PDFs, together with the optimal maximum likelihood decision threshold curve which minimizes the total decision error probability.

refers to the PTEN data of the first and the 10th hours, and $N = 3$ indicates the PTEN data of the first, the 10th, the 20th hours, etc. This assumes at any given time, decision is made based on the data of that given time, plus the data of the previous time instants, which means progressively accumulating the data to make decisions. It is observed in Fig. 9B that P_E first decreases, and after a certain point, it remains nearly constant. To understand this behavior, we note that if the data collected at various time instants are independent, then error probability of a decision-making system that performs sequential hypothesis testing decreases, as the number of observations N increases [42]. This property of a multivariate sequential decision-maker is intuitively appealing. However, if the data collected at various time instants are correlated, the performance of the multivariate sequential decision-maker can significantly degrade, and its error probability does not necessarily decrease, as N increases [42].

To examine possible temporal correlations among the data that the suggested sequential decision strategy employs, we compute condition numbers of Σ_0 and Σ_1 , the $N \times N$ covariance matrices of the data for the two hypotheses H_0 and H_1 , for IR = 1 and 2 Gy, respectively, as N increases from 2 to 8 (Fig. 9C). The condition number of a matrix is the ratio of its largest singular value to its smallest. A large condition number indicates that the matrix is nearly singular. On the other hand, a near singular covariance matrix of several random variables means that some of the random variables are highly correlated. Therefore, a large condition number for a covariance matrix implies large correlations among some of its random variables. We observe in Fig. 9C that as N increases, condition numbers of both of the covariance matrices Σ_0 and Σ_1 increase. This means as time evolves after a certain point, the suggested sequential decision-maker incorporates a new observation that is correlated with the previously used observations. The correlation does not allow

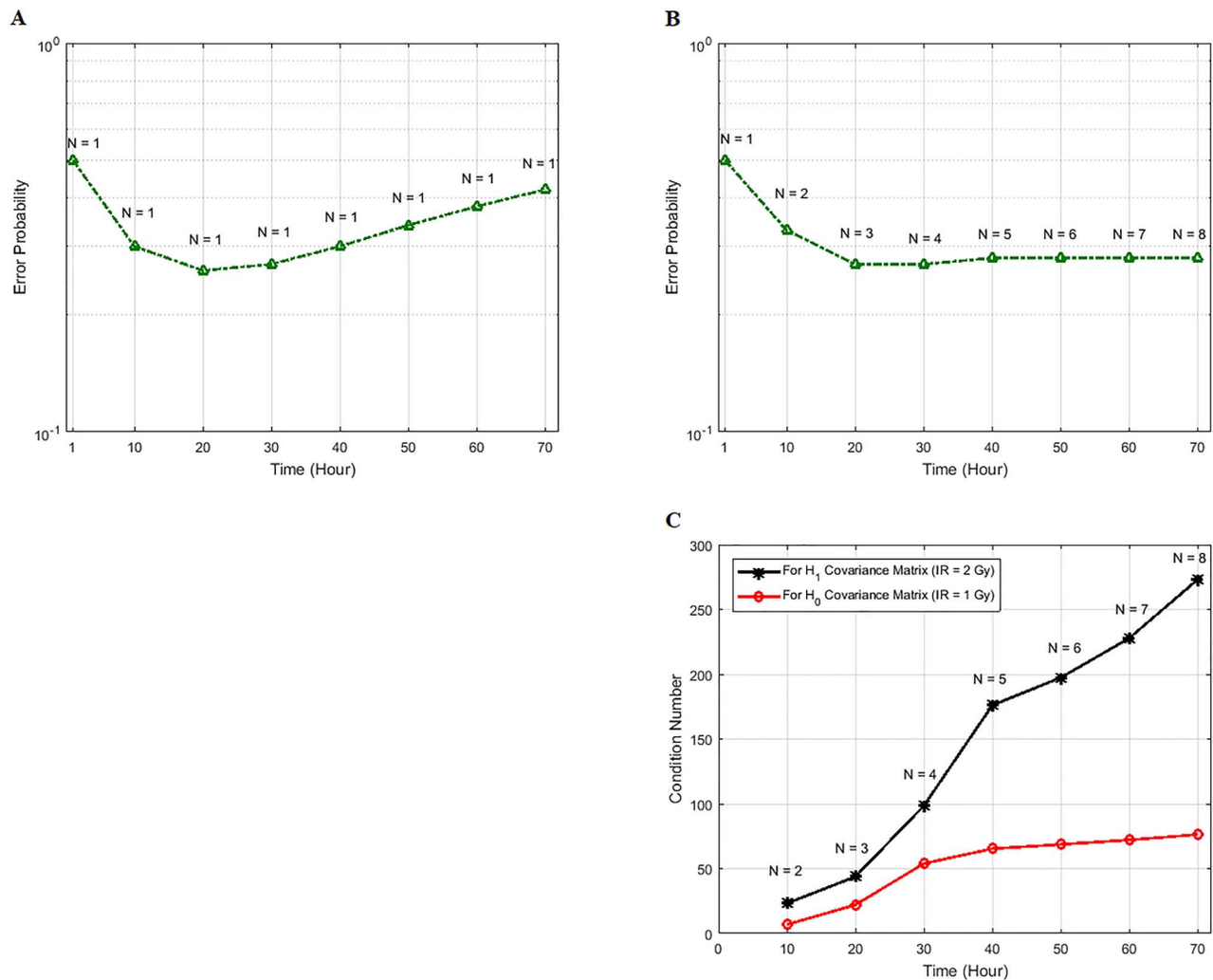


Figure 9. Decision error probabilities versus time in the normal p53 system: a single versus multiple time point study. (A) P_E as a function of time for the 1 vs. 2 Gy radiation scenario, computed using only the PTEN data of a single, $N = 1$, individual time instant. This assumes at any given time, decision is made based on the data of that time only. Having a minimum error probability at the 20th hour is noteworthy. (B) P_E as a function of time for the 1 vs. 2 Gy radiation scenario, computed using the PTEN data of N time instants, $N = 1, 2, \dots, 8$ ($N = 1$ means the PTEN data of the first hour, $N = 2$ refers to the PTEN data of the first and the 10th hours, $N = 3$ indicates the PTEN data of the first, 10th, 20th hours, etc.). This assumes, at any given time, decision is made based on the data of that time, plus the data of the previous time instants, which means accumulating the data to make a decision. It is observed that P_E first decreases, and after a certain point, it remains nearly constant. (C) Condition numbers of Σ_0 and Σ_1 , the $N \times N$ covariance matrices of the data for the two hypotheses H_0 and H_1 , for IR = 1 and 2 Gy, respectively, as N increases from 2 to 8. When N increases, condition numbers of both of the covariance matrices Σ_0 and Σ_1 increase. On the other hand, a large condition number for a covariance matrix implies large correlations among some of its random variables. Therefore, as time evolves after a certain point, the suggested sequential decision-maker incorporates a new observation that is correlated with the previously used observations. The correlation does not allow the decision error probability P_E to decrease beyond a certain point, although N constantly increases.

the decision error probability to decrease beyond a certain point, although N constantly increases (Fig. 9B).

Multivariable analysis of two or more molecules: methods for computing decision thresholds and decision error rates using their concentration measurements in individual cells

So far we have focused on multivariable decision-making and signaling outcome analysis for one molecule at different time instants. However, the introduced methods and algorithms are not limited to the outcome analyses for just one molecule, and they can be applied to various other scenarios and studies. In fact, they can be used to analyze and compute decision error rates based on concentration levels of two or more molecules,

measured simultaneously or even at different time instants. For example, if decision and outcome analysis are going to be conducted based on simultaneous concentration level measurements of two molecules labeled by x and y , then Equations (10–14) can be used to find the maximum likelihood bivariate decision strategy and its minimum error probability. As a more elaborate example, suppose concentration levels of molecule A measured at time instants t_1 and t_2 are labeled as variables x and y , respectively, concentration levels of molecule B measured at t_1 and t_2 are labeled as variables v and w , respectively, and finally concentration levels of molecule C measured at t_1 and t_2 are labeled as variables ψ and ζ , respectively. The 6×1 decision vector ω including all these six decision variables can be defined as $\omega = [x \ y \ v \ w \ \psi \ \zeta]^T$, where T stands for transpose. Now Equations (15–17) can be used to find the maximum likelihood

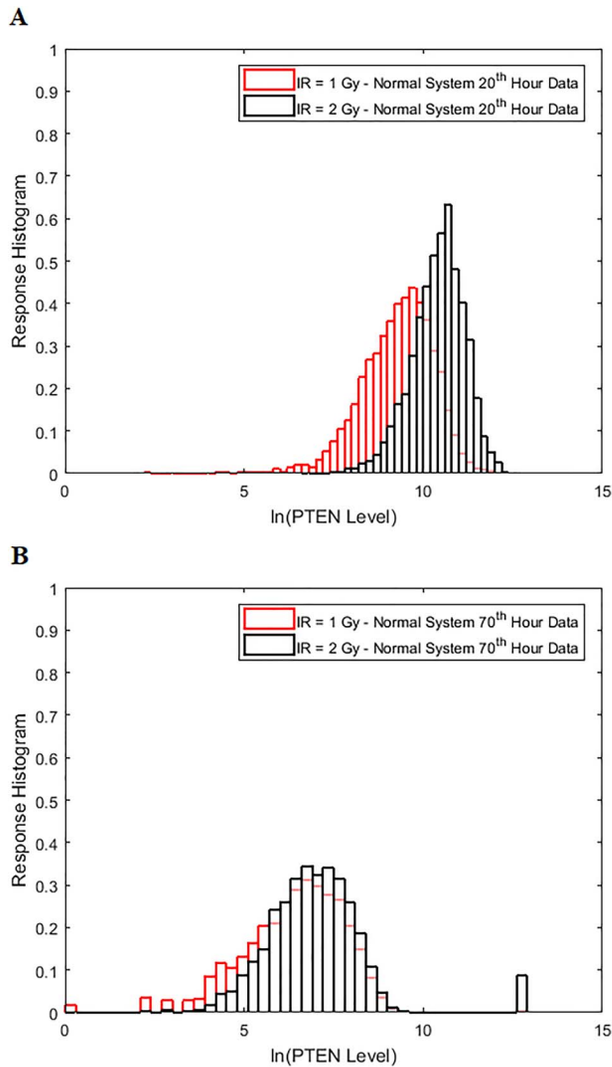


Figure 10. Comparison of the histograms of cell PTEN levels at the 20th and 70th hours under IR = 1 and 2 Gy doses in the normal p53 system. (A) Histograms of the 20th hour PTEN data under IR = 1 and 2 Gy doses, which show less overlap. (B) Histograms of the 70th hour PTEN data under IR = 1 and 2 Gy doses, which show more overlap.

six-variate decision strategy and its minimum decision error probability.

Effect of heterogeneity of initial values and reaction rates on cell response histograms

In addition to stochasticity in dynamic processes, it is natural to consider that the initial level of each protein is not the same in a cell population (heterogeneity of cells). Additionally, there are pseudo-first-order dephosphorylation reactions for which reaction rate coefficients depend on the levels of implicit phosphatases. Thus, it is also natural to assume that reaction rate coefficients corresponding to pseudo-first-order dephosphorylations may vary from cell to cell. To see the effect of heterogeneous initial values and parameters on cell response distributions, we generated new PTEN data for IR = 2 Gy in 5000 cells, assuming that the initial values and parameters of the p53 system are coming from lognormal distributions with means equal to their default values, and ran simulations for different

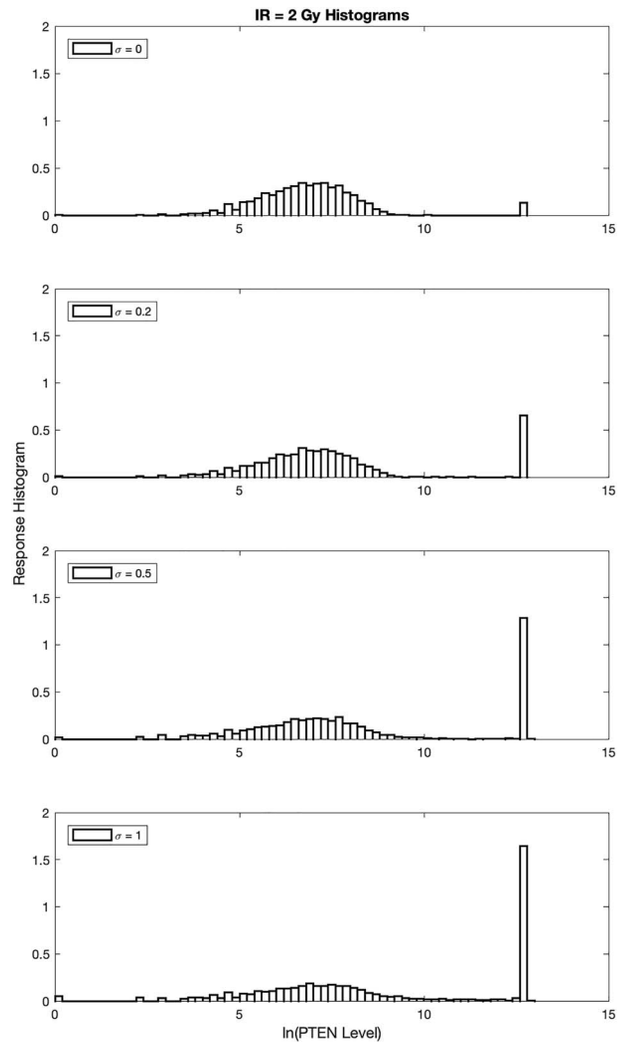


Figure 11. Effect of heterogeneity of initial values and pseudo-first-order dephosphorylation reaction rates on PTEN histograms. Histograms of PTEN levels of cells under IR = 2 Gy dose, with $\sigma = 0, 0.2, 0.5$ and 1.

standard deviations, i.e. $\sigma = 0.2, 0.5$ and 1 [43]. Cell response histograms of the new data for different σ values are shown in Fig. 11 and compared against the 2 Gy data of homogenous cells ($\sigma = 0$). In this system and example, we observe that PTEN histograms undergo some change as σ increases, i.e. more cell heterogeneity, which may result in some changes in decision error probabilities. Nevertheless, one can still use the exact same methods and algorithms introduced in the paper, to conduct signaling outcome analyses of interest for inhomogeneous cells.

BEYOND BINARY DECISIONS: TERNARY DECISIONS AND SIGNALING OUTCOMES AND TERNARY ERROR PROBABILITIES

While the focus of this paper is on binary hypothesis testing, it is possible to develop a multiple hypothesis testing model for outcome analysis, where there exist more than two possible outcomes. This entails more erroneous decisions than false alarm and miss events. Optimal decision thresholds and

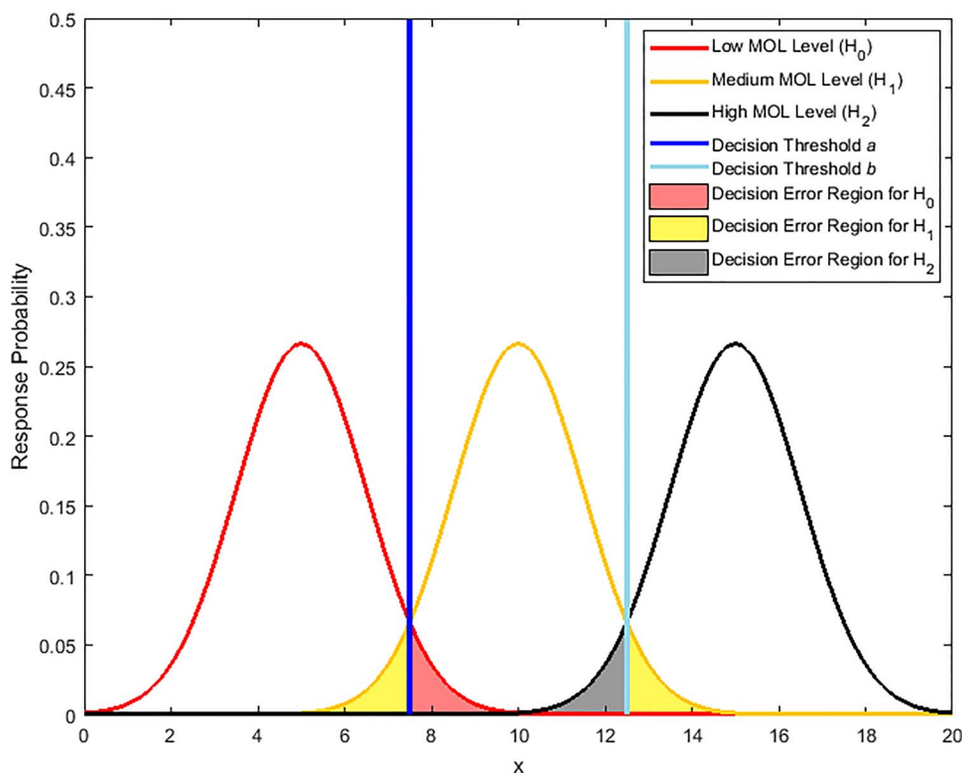


Figure 12. Response PDFs of a hypothetical molecule called MOL whose level entails a ternary decision-making process with three signaling outcomes. Shaded tail areas with the same color represent decision error regions associated with each specific hypothesis. Assuming equi-probable hypotheses, optimal maximum likelihood decision thresholds which minimize the total decision error probability are shown by vertical blue lines at the points of intersection of the probability density functions.

error probabilities for all the incorrect decisions can be similarly computed. For example, assume there are three different signaling outcomes depending on concentration level of a hypothetical molecule called MOL, whose level can fall within one of three regions, which results in the following three possible hypotheses:

$$\begin{aligned} H_0 &: \text{MOL level is low,} \\ H_1 &: \text{MOL level is medium,} \\ H_2 &: \text{MOL level is high.} \end{aligned} \quad (18)$$

Let us assume under each condition, PDF of the MOL level represented by x is normal or Gaussian, i.e. $x \sim \mathcal{N}(\mu_i, \sigma^2)$ such that $\mu_0 < \mu_1 < \mu_2$, where variances are assumed to be equal, to simplify the notation. These PDFs are shown in Fig. 12, with $\mu_0 = 5, \mu_1 = 10, \mu_2 = 15$, and $\sigma^2 = 2.25$. By extending the binary decision errors presented earlier in Equations (8) and (9), ternary decision errors for the three hypotheses can be written as

$$P_{E,H_0} = \int_a^{\infty} p(x|H_0) dx = Q\left(\frac{a - \mu_0}{\sigma}\right), \quad (19)$$

$$P_{E,H_1} = \int_{-\infty}^a p(x|H_1) dx + \int_b^{\infty} p(x|H_1) dx = Q\left(\frac{\mu_1 - a}{\sigma}\right) + Q\left(\frac{b - \mu_1}{\sigma}\right), \quad (20)$$

$$P_{E,H_2} = \int_{-\infty}^b p(x|H_2) dx = Q\left(\frac{\mu_2 - b}{\sigma}\right). \quad (21)$$

In the above equations, a and b are thresholds to decide between H_0 and H_1 and between H_1 and H_2 , respectively. This means the decision regions for the three hypotheses can be written as:

$$\begin{aligned} H_0 &: x < a, \\ H_1 &: a < x < b, \\ H_2 &: b < x. \end{aligned} \quad (22)$$

For equi-probable hypotheses and similar to the derivation that leads to Equation (7), optimal decision thresholds which minimize the total decision error probability can be shown to be

$$a = \frac{\mu_0 + \mu_1}{2}, \quad b = \frac{\mu_1 + \mu_2}{2}. \quad (23)$$

Upon substituting (23) in (19–21), the total error probability in making ternary decisions can be written as:

$$\begin{aligned} P_E &= (1/3)P_{E,H_0} + (1/3)P_{E,H_1} + (1/3)P_{E,H_2} \\ &= (1/3)Q\left(\frac{\mu_1 - \mu_0}{2\sigma}\right) + (1/3)\left[Q\left(\frac{\mu_1 - \mu_0}{2\sigma}\right) + Q\left(\frac{\mu_2 - \mu_1}{2\sigma}\right)\right] \\ &\quad + (1/3)Q\left(\frac{\mu_2 - \mu_1}{2\sigma}\right) \\ &= (2/3)Q\left(\frac{\mu_1 - \mu_0}{2\sigma}\right) + (2/3)Q\left(\frac{\mu_2 - \mu_1}{2\sigma}\right). \end{aligned} \quad (24)$$

As a reference, for the binary decision-making problems and outcome analysis studied earlier in the paper and using Equations (8) and (9), the total error probability in making binary decisions with equal variances simplifies to

$$P_E = (1/2)Q\left(\frac{\mu_1 - \mu_0}{2\sigma}\right) + (1/2)Q\left(\frac{\mu_1 - \mu_0}{2\sigma}\right) = Q\left(\frac{\mu_1 - \mu_0}{2\sigma}\right). \quad (25)$$

To compare ternary and binary error probabilities, let us assume $\mu_2 - \mu_1 = \mu_1 - \mu_0 = \gamma$, which reduces Equations (24) and (25) to $(4/3)Q(\gamma/(2\sigma))$ and $Q(\gamma/(2\sigma))$, respectively. This indicates that the ternary decision error rate can be higher than the binary decision error rate, under the assumed conditions.

ON THE COSTS OF CORRECT AND INCORRECT DECISIONS

In decision theory, there can be some costs associated with correct or incorrect decisions. Let C_{ij} be the cost of deciding H_i when H_j is true. To minimize the expected cost, $C_{00}P(H_0) + C_{01}P(H_1)P_M + C_{10}P(H_0)P_{FA} + C_{11}P(H_1)$, the decision-making system decides H_1 if [30]

$$L(x) = \frac{p(x|H_1)}{p(x|H_0)} > \frac{(C_{10} - C_{00})P(H_0)}{(C_{01} - C_{11})P(H_1)} = \gamma, \quad (26)$$

where $C_{10} > C_{00}$ and $C_{01} > C_{11}$. Usually the costs associated with correct decisions are zero, i.e. $C_{00} = C_{11} = 0$. Additionally, if there is no preference in assigning different costs to different incorrect decisions, one can choose $C_{10} = C_{01}$. This is what we would consider as well, since we do not have a knowledge of the costs of incorrect decisions in the studied cellular system. Upon substituting $C_{00} = C_{11} = 0$ and $C_{10} = C_{01}$ in the above equation, it simplifies to the following equation, which is the optimal maximum likelihood decision rule presented earlier in the paper:

$$L(x) = \frac{p(x|H_1)}{p(x|H_0)} > \frac{P(H_0)}{P(H_1)} = \gamma. \quad (27)$$

CONCLUSION

This study presents a set of decision-theoretic, statistical signal processing and machine learning methods and metrics for modeling and measurement of decision-making processes and signaling outcomes under normal and abnormal conditions and in the presence of noise and other uncertainties. Due to the noise, signaling malfunctions or other factors, cells may respond differently to the same input signal. Some of these responses can be erroneous and unexpected. Here we present univariate and multivariate models and methods for decision-making processes and signaling outcome analyses and, as an example, apply them to an important system that is involved in cell survival and death, i.e. the p53 system shown in Fig. 1 (another decision analysis example can be found in the paper by Habibi et al. [1]). The p53 system becomes active due to DNA damage caused by IR, and as a result, a cell can take two different actions: it can either survive by repairing the DNA or trigger apoptosis. In this context, we model decisions and signaling outcomes triggered by the p53 system as a binary hypothesis testing problem, where two hypotheses are introduced in Equation (1). Regarding these two hypotheses, our approach identifies that there can be two types of incorrect decisions: ‘false alarm’ and ‘miss’. To compute the likelihood of these decisions, we employ the simulator of Hat et al. [9], to obtain single cell data of the p53 system, by exposing the cells to different radiation doses. We consider PTEN levels in cells as the decision variable, since it is a good predictor of cell fate [9]. Our analysis focuses on low radiation dose versus high radiation dose scenarios, where we fix the low IR dose at 1 Gy, whereas we set the high IR dose at 2–8 Gy. We also analyze decision-making events and signaling outcomes when an abnormality is present in the p53 system.

The incorrect decision probabilities provided in Equation (2) and the overall decision error probability in Equation (3) are computed after determining an optimal decision threshold. We obtain this decision threshold using the maximum likelihood principle which states that the best decision can be made by selecting the hypothesis that has the maximum probability of occurrence. We compute decision threshold and error probabilities using single time point data of PTEN levels in both normal and abnormal p53 systems. For 1 vs. 2 Gy and 1 vs. 8 Gy case studies, we present histograms, response distributions, decision thresholds and false alarm and miss decision regions in normal and abnormal p53 systems in Figs. 3 and 5, respectively. Our decision analysis reveals and quantifies that more erroneous decisions are made when deciding between two nearly the same radiation doses in the normal p53 system (Fig. 6). On the other hand, the difference between responses is easily identifiable for very low versus very high IR doses. This feature seems not to be present in the abnormal p53 systems (Fig. 6), according to our decision modeling approach. Our decision and outcome analyses and observations are further visualized and confirmed by using the ROC curves (Fig. 7), which are useful graphical tools to study the performance of decision-making systems. We would like to note that these observations are specifically made based on the low versus high IR case studies, e.g. d_0 vs. d_1 IRs introduced in the paper for the p53 system, as an example of a signaling network, in which the low IR dose is fixed to 1 Gy ($d_0 = 1$ Gy) and the high IR dose is ranging from 2 up to 8 Gy ($d_1 = 2, 3, \dots, 8$ Gy). Such conclusions may not be generalized to other biological hypotheses and systems, while the proposed framework and its analytical tools, whose introduction has been the main goal of this paper, can be similarly used.

In addition to the above univariate single time point analysis, we extend our signaling outcome modeling framework to dynamical multi-time point measurements and multi-dimensional decision making algorithms, to see how the number of decision variables affects the decisions and signaling outcomes over time. To introduce the concepts, first we conduct a bivariate analysis, for which bivariate response distributions of cell PTEN levels measured at two different time instants are shown in Fig. 8, as well as the optimal maximum likelihood decision boundary. Then we introduce a multivariate dynamic decision modeling framework, for the general scenario where there are more than two decision variables over time. This allows to model and understand how decision error probability changes over time, if at any time the decision is made based on the current observation, together with the previous observations. We observe in Fig. 9B that as the decision-making strategy incorporates more and more PTEN data of various time instants into its decisions, for the p53 system exposed to two radiation doses of 1 and 2 Gy, the decision error probability reaches its smallest value at a certain time instant. However, adding more data afterwards does not necessarily improve the decision precision, i.e. the decision error probability does not necessarily decrease as N increases with time (Fig. 9B). We show that this behavior can be related to the correlations that exist among the PTEN levels measured at different times (Fig. 9C).

Although we focus on multivariable decision-making and signaling outcome analysis for one molecule at different time instants, the introduced methods and algorithms are not limited to the outcome analyses for just one molecule. They can be applied to various other scenarios and studies. For instance, they can be used to analyze decision strategies and compute decision error rates based on concentration levels of two or more

molecules, measured simultaneously or even at different time instants.

We finally show how the introduced binary decision-making and signaling outcome analysis models can be extended to more than two decisions, i.e. more than two hypotheses. A ternary scenario with three signaling outcomes is analyzed as an example, and it is shown that under certain conditions, ternary decision error probability can be higher than the binary one.

The methods and models presented here can be expanded to describe the performance and precision of more complex systems and networks such as the ones whose inputs are multiple ligands or secondary messengers and whose outputs are several transcription factors involved in certain cellular functions. Analyzing concentration levels of these transcription factors over time using the proposed approaches can model various decisions and signaling outcomes, and their probabilities, in the presence of noise or some cellular abnormalities and in response to the input signals.

The methods and formalism developed in this study are applicable to a wide variety of signaling outcome analyses, decision-makings and signal transduction processes where there are two or more possible outcomes. For example, in the context of *Escherichia coli* chemotaxis, binary decisions (influencing all chemotaxis processes) are either to continue motion in the same direction or to change the flagellum operation mode from run, counterclockwise, to tumble, clockwise, resulting in random direction changes [44]. Based on the network or system of interest and the available data, the hypotheses in Equation (1) can be revised, and subsequently the same mathematical framework and algorithms and methods can be applied, using the underlying probability distributions of data.

Overall, these decision-theoretic models and signaling outcome analysis methods can be beneficial for better understanding of transition from physiological to pathological conditions such as inflammatory diseases, various cancers and autoimmune diseases.

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CONFLICT OF INTEREST STATEMENT

The authors declare that no conflicts of interest exist.

AUTHOR CONTRIBUTIONS

M.O. developed the methods, performed the simulations and computations and wrote the manuscript. T.L. contributed to and discussed the research topics and edited the manuscript. A.L. contributed to and discussed the research topics and edited the manuscript. E.S.E. contributed to and discussed the research topics and edited the manuscript. A.A. conceived the ideas and concepts, developed the methods and wrote the manuscript.

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