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Median Effect Dose and Combination Index Analysis of Cytotoxic Drugs Using Flow Cytometry

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1. Introduction

Targeted therapy is a strategy of anticancer treatment that aims to interfere with processes of tumorigenesis, cancer progression and metastasis by selectively affecting key molecules of tumor cells (Armand et al., 2007; Favoni & Florio, 2011; Gross-Goupil & Escudier, 2010). Targeted therapies are directed to small molecules participating in different mechanisms that control cell survival through cellular proteins or signalling pathways (Mueller et al., 2009; Zahorowska et al., 2009). Targeted therapies may offer enhanced efficacy, enhanced selectivity, and less toxicity. However, targeting selective molecules and pathways often induces the activation of redundant mechanisms and enhances the emergence of resistant cells due to selective pressure (Woodcock et al., 2011). This is one of the reasons why the effects of targeted agents are not durable when used alone, and often result in drug resistance and clinical relapse.

Except for specific cases the use of these targeted drugs as monotherapy is often discouraged due to lack of efficacy. However, combined therapy with drugs targeting several mechanisms of tumor cell death can greatly improve efficacy and may overcome resistance. Several genomic and epigenetic alterations have been identified in tumor cells that lead to unrestrained proliferation, evasion of proapoptotic signals, metastasis, and resistance to drug-induced cell death. These alterations are critical for cancer progression and therefore combination strategies employing multiple targeted agents can be a successful therapeutic strategy. In vertical combination strategies two or more drugs target a same pathway at two different points, while in horizontal combinations drugs are directed towards different intracellular signalling pathways and have the potential advantage of combining agents with non-overlapping toxicities (Gross-Goupil & Escudier, 2010).

Novel treatments require the investigation of mechanisms of action and synergy of combination treatments to enhance the role of the targeted pharmacological agents (Carew et al., 2008; Mitsiades et al., 2011). Evaluating combinations of targeted drugs, including investigational agents, are an essential part of this effort (Dancey & Chen, 2006). An interesting example is represented by Bortezomib, a drug currently effective as single agent in multiple myeloma and mantle cell lymphoma (Bross et al., 2004; Kane et al., 2007; Wright, 2004).
Bortezomib is a proteasome inhibitor that selectively triggers apoptosis in various types of neoplastic cells. It has been tested in a wide variety of solid tumors but has generally been ineffective as monotherapy (Boccadoro et al., 2005). However, Bortezomib has shown increased activity when combined with several novel targeted agents including protease inhibitors, kinase inhibitors, farnesyltransferase inhibitors, HSP-90 inhibitors, pan-Bcl-2 family inhibitors, and other classes of targeted inhibitors (Dai et al., 2003; Karp & Lancet, 2005; Pei et al., 2004; Perez-Galan et al., 2008; Trudel et al., 2007; Workman et al., 2007; Yanamandra et al., 2006). Thus, Bortezomib in combination with novel targeted therapies increase antitumor activity and overcome specific cellular antiapoptotic mechanisms (Wright, 2010).

Two-drug combination therapies are being assessed in a variety of tumors, usually testing agents that have different targets, nonoverlapping toxicity, and some rationale for evaluation (Belinsky et al., 2011; Castaneda & Gomez, 2009; Eriksen et al., 2009; Klosowska-Wardega et al., 2010). An increase in the number of these studies is expected in coming years, on the basis of emerging data with new agents, which is expanding our understanding of the molecular pathways important in cancer progression. (Woodcock et al., 2011). Tumor intracellular signalling pathway dependencies are being increasingly analyzed, and patients treated on the basis of the resistance profile detected for specific drug combinations (Busch et al.;Derenzini et al., 2009; Michiels et al., 2011). This approach may facilitate the development of combination regimens optimized for specific tumor subtypes, thus providing the potential for tailored therapy in individual patients on the basis of certain molecular and genetic characteristics of their disease.

1.1 Targeted drugs often induce programmed cell death as their main mechanism of anti-tumor activity

Many of the classic chemotherapeutic agents (alkylating agents, antimetabolites, antibiotics, topoisomerase inhibitors) are known to block cell division by compromising DNA replication and halting cell cycle progression, or inhibit mitosis, eventually leading to cell death (Foye, 1995; Goodman et al., 2010; Shuck & Turchi, 2010). Indicators of cell proliferation are suitable effect biomarkers to assess whether a combination of these agents is synergic, additive or antagonistic. Biomarkers frequently used for this purpose include incorporation of nucleotide analogues such as bromodeoxyuridine, or metabolic indicators of cell number such as tetrazolium salt-based assays (Karaca et al., 2009; Olszewska-Slonina et al., 2004; Sims & Plattner, 2009). However many of the new targeted agents interfere with constitutively active survival pathways or initiate apoptosis by directly influencing pro-apoptotic signals (Citri et al., 2004; Kim et al., 2005; Larsen et al., 2011; Vega et al., 2009; Zhang et al., 2009). In addition autophagic cell death and programmed necrosis are being actively investigated as alternative and pharmacologically relevant forms of programmed cell death (Berghe et al., 2010; Bijnsdorp et al., 2011; Duan et al., 2010; Gozuacik & Kimchi, 2007; McCall, 2010; Notte et al., 2011; Paglin et al., 2005; Platini et al., 2010). Combination studies should be conducted using effect biomarkers that are as close as possible to the known mechanisms targeted by single agents, and biomarkers specifically related to drug-induce tumor cell death appear more adequate for the assessment of new targeted agents (Cameron et al., 2001; Facoetti et al., 2008; Wesierska-Gadek et al., 2005). A straightforward approach is to determine the proportion of live and dead cells in viability studies scoring
thousands of cells through flow cytometry which ensures exceptional precision for dose-effect cytotoxicity studies.

2. Assessment of viability through flow cytometry

The strength of flow cytometry when compared to other methods available to assess the proportion of live and dead cells is the accuracy and precision brought by single cell multiparametric assessment. A variety of fluorescent probes may be chosen to use in viability assessment through flow cytometry. These probes are based on cell functions and biological conditions that are differentially preserved in live cells and lost in dead cells. It is usual to select at least two probes measuring independent functional conditions. For example, a probe evaluating membrane integrity of cells and another probe evaluating enzymatic activity. Probes should be selected to match specific experimental requirements such as biological variability, duration of the experiments, whether cells exposed to drugs are adherent or non-adherent, and illumination lines available in the flow cytometer. Some probes may be released after being retained within the cells for a short time and require immediate assessment through the flow cytometer after labelling, while others may be retained for several hours or may be even retained indefinitely by being covalently linked to cellular components. In addition some specimens may require fixation due to biohazard issues, so another kind of probes should be chosen and combined in these cases (De Clerck et al., 1994).

2.1 Fluorescent staining of live and dead cells

Viability is not easily defined in terms of a single physiological or morphological parameter. No single parameter fully defines cell death in all systems; therefore, it is often advantageous to use more than one cell death indicators based on different parameters such as membrane damage, and enzymatic or metabolic activity. A considerably large number of fluorescent probes have been introduced in the recent years that are dedicated to the assessment of viability on a single cell basis. Many of these new probes have features that are useful under specific experimental circumstances. The two conditions most often detected are increased cell membrane permeability in dead cells and the presence of enzymatic activity in live cells. The former is assessed with probes that become fluorescent when bound to DNA but are not able to pass through cell membrane if selective permeability is preserved, while the later is determined by fluorogenic substrates. However other conditions occurring only in live cells may be used to demonstrate viability such as enzymatic oxidation, reduction and mitochondrial membrane potential (Callewaert et al., 1991). It is important to underscore this concept because these probes are often used for assessment of specific cellular functions and it may be erroneously assumed that they have no contribution to the assessment of viability.

2.2 Enzymatic activity in live cells, use of tracker dyes

One of the first probes introduced and most frequently used to stain live cells has been fluorescein-diacetate (FDA) (Jones & Senft, 1985; Ross et al., 1989). This non-fluorescent cell-permeant esterase substrate penetrates the cell and is converted by nonspecific intracellular esterases into fluorescein.
Thus it becomes a more polar compound and is retained within those cells that have intact plasma membranes. In contrast, nonfluorescent FDA and fluorescein rapidly leak from those cells that have a damaged cell membrane because it is no longer retained due to increased permeability (Prosperi et al., 1986). This property ensures that dead cells will never retain FDA or fluorescein, even if cell death occurs after the labelling procedure. This is one of the reasons why it is recommended to analyze cells rapidly through the flow cytometer after staining with FDA and why they should be kept in low incubation temperatures to minimize potential fluorescein leakage.

Calcein-acetoxyethyl-ester (Calcein-AM) is a derivative of calcein that has several improvements over FDA (Duan et al., 2010; Papadopoulos et al., 1994). Calcein-AM is also a substrate of nonspecific intracellular esterases. The fluorescent product is calcein and is better retained in cells because it is a polyanionic compound that has six negative charges and two positive charges at pH 7. Calcein-Blue-AM and Calcein-Violet-AM are similar to Calcein-AM but have different excitation and emission wavelengths (Fuchs et al., 2007). Calcein-Blue-AM is excited with UV lasers while Calcein-Violet-AM is excited with 405 nm violet diode lasers, although both dyes emit blue fluorescence (Prowse et al., 2009). They can be used when the green fluorescence channel from the 488 nm excitation line is needed for other purpose and a UV or violet illumination line is available. Chloromethyl-fluoresceindiacetate (CM-FDA), is a FDA derivative that is retained within the cell even after damage to the plasma membrane due to its ability to bind thiol groups (Lantz et al., 2001; Sarkar et al., 2009). The weakly thiol-reactive chloromethyl moieties of this compound react with intracellular thiols and the acetate groups are cleaved by cytoplasmic esterases (West et al., 2001). This compound will not stain dead cells but the label will be preserved in those cells that die after the labelling procedure because the fluorescent product will be bound to SH groups within the cells (Sebastia et al., 2003). Chloromethyl SNARF-1 acetate is similar to CM-FDA but exhibits red fluorescence when excited with 488 nm blue laser. Thus it can be used when the green fluorescence channel is needed for other purpose and a UV or violet illumination line is not available (Hamilton et al., 2007). Carboxi-fluorescein-succinimidyl-ester (CFSE) is converted to fluorescent compound by intracellular esterases but covalently

| Probe            | Excitation line | Fluorescence emission | Intracellular retention |
|------------------|-----------------|-----------------------|-------------------------|
| FDA              | Blue            | Green                 | Poor                    |
| Calcein-AM       | Blue            | Green                 | Good                    |
| Calcein Blue-AM  | UV              | Blue                  | Good                    |
| Calcein Violet-AM| Violet          | Blue                  | Good                    |
| 5-Cl-M-FDA       | Blue            | Green                 | Excellent               |
| 5-Cl-M-SNARF     | Blue            | Orange                | Excellent               |
| CFSE             | Blue            | Green                 | Excellent               |

Table 1. Fluorogenic substrates of intracellular esterases that are commonly used as viability probes
binds amino groups of proteins and is completely retained within cells, even after damage of cell membrane (Fujioka et al., 1994; Li et al., 2003). This dye is also used as cell tracker because it is retained in daughter cells after several rounds of cell division (Parish & Warren, 2002). It is worth to note that probes like FDA may give poor results with trypsinized cells owing to potential leakage of fluorescein during the staining and washing procedures (Zamai et al., 2001). Thus probes like CM-FDA, CM-SNARF-1, and CFSE may be a better choice for staining adherent cells.

2.3 DNA labelling in live and dead cells

Many polar nucleic acid stains are able to enter eukaryotic cells only when the plasma membrane is damaged. These stains are known as cell-impermeant dyes and include propidium iodide (PI) which is the most frequently used probe for assessing viability in flow cytometry (Yeh et al., 1981). This dye is excluded from live cells because it is negatively charged but readily penetrates the membrane of damaged cells and binds DNA. When excited at 488 nm DNA-bound PI increases orange-red fluorescence emission more than 1000 fold. Another commonly used cell-impermeant dye excited with 488 nm laser is 7-aminoactinomycin-D (7AAD). This dye binds DNA only in dead cells but emits fluorescence beyond 610 nm and allows the usage of the yellow-orange fluorescence channel for other purpose (Pallis et al., 1999).

Both PI and 7AAD have large Stokes shifts and can be used in 488 nm laser flow cytometers with green fluorescent tracker dyes such as FDA, CM-FDA, Calcein-AM and CFSE. Cells with damaged membranes may be identified with other cell-impermeant DNA fluorescent dyes that emit fluorescence in different wavelengths than that of PI.

The SYTOX series includes SYTOX-green (excited with 488 nm laser), SYTOX-red (excited with 633 and 635 nm lasers) and SYTOX-blue (excited with UV or 405 nm violet diode laser) (Haase, 2004; Lebaron et al., 1998; Yan et al., 2005). In contrast to SYTOX dyes, the SYTO series of nucleic acid stains can enter live cells and are thus cell-permeant DNA dyes (Ullal et al., 2010). The SYTO probes bind DNA with low affinity in live or dead cells (Eray et al., 2001; Poot et al., 1997). They are combined with high affinity cell-impermeant dyes to discriminate live from dead cells (Wlodkowic & Skommer, 2007). For example cell-permeant SYTO-red will stain live and dead cells with red fluorescence binding with low affinity to DNA, but if used together with SYTOX-green dead cells will be green fluorescent, because SYTOX-green has much higher affinity for DNA and will displace the low affinity SYTO-red. In addition, SYTOX-green will never stain live cells because it is cell-impermeant.

2.4 Biohazardous specimens

Viability staining of biohazardous specimens often requires fixation procedures that inactivate pathogens but produce minimal distortion of cellular characteristics. Some combinations of cell permeant and cell impermeant DNA dyes can be treated with fixatives such as 4% glutaraldehyde or formaldehyde to allow safer handling during analysis, without disrupting the distinctive staining pattern. An example is provided by cell-permeant, green-fluorescent DNA probe SYTO-10 and the cell-impermeant, red-fluorescent DNA probe ethidium homodimer-2 (Barnett et al., 2004; Poole et al., 1996). Using these two probes cells can be stained and fixed at various times during an experiment, and the results
can be analyzed several hours later. This method may be applied to viability assessment of any non-adherent cells, as well as trypsinized adherent cells. Tables 1, 2, and 4 summarize the main features of viability probes based on enzymatic activity and DNA labelling discussed above that may be considered to meet specific experimental requirements.

| Probe              | Excitation line | Fluorescence emission | Membrane Permeant | DNA Affinity | Fixable |
|--------------------|-----------------|-----------------------|--------------------|--------------|---------|
| SYTOX-Blue         | UV-Violet       | Blue                  | NO                 | High         | NO      |
| SYTOX-Green        | Blue            | Green                 | NO                 | High         | NO      |
| SYTOX-Red          | Red             | Red                   | NO                 | High         | NO      |
| Propidium iodide   | Blue            | Orange-Red            | NO                 | High         | NO      |
| 7AAD               | Blue            | Red                   | NO                 | High         | NO      |
| SYTO-10            | Blue            | Green                 | YES                | Low          | YES     |
| SYTO-Red           | Blue            | Red                   | YES                | Low          | NO      |
| Ethidium homodimer-2 | Blue         | Green                 | NO                 | High         | YES     |

Table 2. DNA probes used for viability assessment

2.5 Two parameter assessment of viability through flow cytometry

Identification of live and dead cells is often conducted with simultaneous use of two probes. The combination may include a cell-impermeant DNA probe and either a fluorogenic substrate or a cell-permeant DNA probe. It should be highlighted that viability may be also indicated by probes that have been designed to assess other cellular functions. For example, generation of hydrogen peroxide and superoxide anion occurs in live cells due to normal function of mitochondrial electron transport chain and does not occur in dead cells. The superoxide anion probe dihydroethidine (HE) and the hydrogen-peroxide probe dihydro-dichloro-fluoresceindiacetate (DH-DCFDA) will stain live cells red fluorescent and green fluorescent respectively (Eruslanov & Kusmartsev, 2010; Zanetti et al., 2005). Both probes may be appropriately combined with cell-impermeant DNA dyes to discriminate between live and dead cells.

Similarly, potentiometric dyes stain live cells with preserved mitochondrial membrane potential, but not dead or compromised live cells where the mitochondrial membrane potential has collapsed (Marchetti et al., 2004). Thus they may also be combined with cell-impermeant DNA dyes to discriminate live and dead cells. For example, rhodamine 123 has been used in combination with propidium iodide for viability assessment with two-color flow cytometry (Darzynkiewicz et al., 1982). Metabolically active cells undergo normal oxidation-reduction reactions and thus can also reduce a variety of probes, providing a measure of cell viability and overall cell health (Callewaert et al., 1991; Radcliff et al., 1991). Resazurin and dodecylresazurin (C12-resazurin) have been extensively used as oxidation-reduction indicators to detect viable cells (Czekanska, 2011). Reduction of resazurin yields
the red fluorescent product resorufin while C12-resazurin yields C12-resorufin which is better retained by single cells (Talbot et al., 2008).

Again these probes may be combined with cell-impermeant DNA probes like SYTOX-green to discriminate live and dead cells with two color flow cytometry.

2.6 Viability assessment with single-color fixable dyes

In some experimental circumstances only one fluorescence channel may be dedicated to assessment of cell viability. In this case amine-reactive fluorescent dyes can be used to evaluate mammalian cell viability.

In cells with compromised membranes, these dyes react with free amines both in the cell interior and on the cell surface, yielding intense fluorescent staining. In viable cells, the dyes only stain cell-surface amines, resulting in less intense fluorescence (Elrefaei et al., 2008). The difference in intensity between the live and dead cell populations is preserved following formaldehyde fixation, using conditions that inactivate pathogens (Burmeister et al., 2008). There are several options of fluorescence excitation (UV, violet, blue, and red lasers) and emission wavelength (blue, green, yellow, red).

| Probe | Excitation line | Fluorescence emission |
|-------|----------------|-----------------------|
| LIVE/DEAD® fixable Blue | UV | Blue-450 nm |
| LIVE/DEAD ® Fixable Aqua | UV | Green-526 nm |
| LIVE/DEAD ® Fixable Yellow | Violet | Yellow-575 nm |
| LIVE/DEAD ® Fixable Violet | Violet | Blue-450 nm |
| eFluor® 450 | Violet | Blue-450 nm |
| Fixable Viability Stain 450 ® | Violet | Blue-450 nm |
| LIVE/DEAD ® Fixable Green | Blue | Green-520 nm |
| eFluor® 506 | Blue | Green-506 nm |
| LIVE/DEAD ® Fixable Red | Red | Red-615 nm |
| LIVE/DEAD ® Fixable Far Red | Red | Far Red-665 nm |
| eFluor® 660 | Red | Far Red-660 nm |

Table 3. Fixable amine-reactive fluorescent probes used for single-color assessment of cell viability. The wavelengths indicated correspond to the emission peaks as specified by the probe manufacturer.
2.7 Viability vs. apoptosis

Most targeted cytotoxic drugs have been shown to induce apoptosis or other modes of programmed cell death, including autophagic cell death or programmed necrosis. These mechanisms of cell death are often contrasted to necrosis where a passive, sudden and uncontrolled disintegration of the cell occurs. Physiological consequences of apoptosis and passive necrosis are different, and thus it is important to determine the cell death phenotype. When assessed through flow cytometry, cells undergoing apoptosis or other forms of programmed cell death show a decrease in cell volume and forward light scatter (FSC), and an increase in side light scatter (SSC) mainly due to cytoplasmic and nuclear changes such as blebbing, and nuclear fragmentations (Dive et al., 1992; Pheng et al., 2000). In contrast necrosis often shows increased cell volume and FSC without changes in SSC (Healy et al., 1998). Viability assessment after cytotoxic drug exposure does not address the cell death phenotype, thus any kind of cell death phenotype may be induce by drug treatment including passive necrosis (Healy et al., 1998). However studies determining the median cytotoxic dose will require exposure to increasing doses from sub-lethal levels to the minimal doses approaching 100% cell death. In this scenario, programmed cell death phenotypes are more frequently observed than passive necrosis.

| Probe   | Membrane Permeant | Excitation line | Fluorescence emission |
|---------|-------------------|----------------|---------------------|
| YOYO-1  | NO                | Blue           | Green               |
| TOTO    | NO                | Blue           | Green               |
| TO-PRO  | NO                | Blue           | Green               |
| POPO-1  | NO                | UV-Violet      | Blue                |
| BOBO-1  | NO                | UV-Violet      | Blue                |
| YOYO-3  | NO                | Red            | Red                 |
| TOTO-3  | NO                | Red            | Far Red             |
| BOBO-3  | NO                | Red            | Red                 |
| JOJO-1  | NO                | Green          | Orange              |
| JO-PRO-1| NO                | Green          | Orange              |

Table 4. Membrane–impermeant dimeric and monomeric cyanine dyes are nonfluorescent unless bound to nucleic acids and have extinction coefficients 10–20 times greater than that of DNA-bound propidium iodide

2.8 FDA-PI staining and the "cell death pathway": Frequency distributions of graded and abrupt transitions

When two fluorescent probes are used to determine the proportion of live and dead cells after exposure to cytotoxic drugs over an extended dose range data analysis would be better analyzed on a bivariate plot.

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An example is the pair represented by FDA as an indicator of esterase activity in live cells and PI as an indicator of cell membrane damage (Fig. 1). In this case a bivariate plot of green fluorescence against red fluorescence will aid in determining the percentage of live and dead cells (Fig. 1A,C). In addition, the bivariate plot will provide useful data about the biological processes evaluated with FDA and PI.

A concept frequently present in flow cytometry, particularly when analyzing bivariate plots, is the presence of graded transitions or abrupt changes (Shapiro, 2003). These patterns in bivariate distributions are determined by the underlying biological process that is being studied. For example damage of cell membrane allows staining by PI probe so that cells may be classified as dead with a permeable membrane or live having preserved selective permeability, depending on whether they are red fluorescent or not. Cells are observed to be bright stained or having no stain at all, but very rarely they are observed to have dim red fluorescence. Thus membrane damage and PI staining is an example of an abrupt transition or discrete process represented by membrane damage that produces a sudden change in the frequency distribution. This frequency distribution is symmetric, bell-shaped, and has low variability around the peak (Fig. 1C,E).

By contrast when analyzing drug-induced effects on esterase activity through green fluorescence we will observe a graded transition from bright fluorescence to dim fluorescence (Fig. 1C). Thus a graded biological process represented by progressively decreased esterase activity determines a skewed frequency distribution with higher variability around the peak (Fig. 1F). In this case there will be a higher probability of observing cells within any level of metabolic activity represented by the amount of green fluorescence: bright, intermediate and dim. Note also that cells with damaged membrane no longer retain fluorescein (very few events are seen in upper right quadrant, Fig. 1C).

When analysis is restricted to live cells without damaged membrane (PI negative) it is more evident that the probability of finding live cells with low metabolic activity in the drug-treated population decreases gradually (Fig. 1H). By contrast, when the analysis is restricted to cells without metabolic activity a narrow bell-shape distribution is observed meaning that the probability of finding cells with damaged membrane in cells without metabolic activity increases abruptly (Fig. 1G). When combined in a bivariate plot the gradual decrease in metabolic activity in live cells is observed as a continuous distribution or pathway, while the abrupt transition from membrane-impermeable to membrane-permeable is observed as a discrete transition to a main single cluster of PI-positive cells with very low or no metabolic activity (Fig. 1C). The probability of finding cells with low or no metabolic activity is very low as shown by the few cells in an intermediate position along this "death-pathway". Changes in FSC and SSC are also graded transitions and define a "death-pathway" in bivariate plots (Fig. 1D). Most cells having membrane damage have low FSC and high SSC, those cells without membrane damage and with metabolic activity have high FSC and low SSC, while intermediate positions may be occupied by either of these populations (Fig. 1D). Thus the death pathway defines a whole range of changes occurring in all four parameter FSC, SSC, FDA, and PI. However the main result is characterizing cells as either dead or alive and this difference is brought by PI staining and membrane damage. Thus applying quadrant analysis we would add the fraction of cells in both upper quadrants and the fraction of cells in the lower quadrants as live cells (Fig 1C). The remaining parameters will work as internal quality controls.
Fig. 1. (continues on next page) The FDA vs. PI bivariate plot and the cell death pathway. A. Sample of human U937 leukemic cells labelled with FDA and PI. Lower right quadrant shows that most cells (97.30%) have esterase enzymatic activity and preserved membrane permeability because they exclude PI staining. Only 1.43% of cells have PI staining without FDA fluorescence, while 1.05% are double positive indicating both enzymatic activity and damaged membrane. B. FSC-SSC profile of live cells is show in green and corresponds to the 97.30% of cells shown in the lower right quadrant of panel A. The small amount of single PI positive (red) and double positive cells (blue) are also observed. C. Sample of human U937 cells exposed to 5 μM sodium arsenite (AsNaO2) for 72h stained with FDA and PI showing a "slow" transition from high FDA fluorescence to low FDA fluorescence (green) and a further "abrupt" transition to a PI positive FDA negative cluster of dead cells (red). A minority of cells are double positive (blue). The whole transition is indicated with the black arrow. D. FSC vs. SSC plot of the sample shown in C. Green color represents FSC-SSC paired values only occupied by live cells (lower right quadrant shown in C), red color represents FSC-SSC paired values only occupied by dead cells (upper left quadrant in C), while yellow color represents FSC-SSC paired values occupied by both live and dead cells. The black arrow shows the graphical death pathway transition in the FSC SSC plot. The FSC-SSC values of the minority of double positive cells are shown in blue.
3. Building a cytotoxic dose response curve

Theoretically, if a population of cells were homogenously sensitive to cell death induced by a certain drug there would be a single dose D at which 100% cell death would be observed (Casarett et al., 2008; Goodman et al., 2010). However in any given sample of drug-treated cells a random proportion of cells will die at doses lower or higher than D due to experimental and biological variability. This random divergence from D follows a Gaussian distribution (Fig. 2A,C).
Fig. 2. Quantal dose response model. A. Single cells will show differences on the minimal drug dose required to induce cell death. Dm is the most frequent minimal dose required to elicit cell death. Variations around this value have a normal distribution. In this particular example Dm is 7uM and the standard deviation (SD) is 1uM. B. Dm is the drug dose that is estimated to kill half of the cells in a sample. When running experiments exposing cells to incremental doses of a cytotoxic drug the fraction of dead cells observed will follow a normal cumulative function. C. Differences between two cytotoxic drugs in the variability observed around Dm. The drug represented by the dotted line has larger variability (SD=1.8 uM) than the drug represented in full line (SD=1 uM). D. Increased variability around Dm is observed as a dose response curve with a smaller slope as shown by the dotted line of the cumulative normal distribution.

The median dose Dm represents a dose D where half of the cells are killed and half of the cells remain live (Fig. 2 B). However, as indicated by the bell shape of the Gaussian distribution many of the 50% of cells killed at a dose Dm may have required less than Dm to be killed. In fact as shown in figure 2A only a fraction of cells will require strictly a dose Dm (those in the bell peak), while a minority of cells will require a dose much lower than Dm to be killed (those in the left tail of the bell-shaped curve). If we conduct experiments evaluating the cytotoxic effect of increasing doses we will observe a sigmoid curve that follows a cumulative Gaussian frequency distribution (Fig. 2B). Doses lower than Dm will show decreased probability of cell death approaching 0% while doses higher than Dm will show increased probability of cell death approaching 100% (Fig. 2B). This model is known as quantal dose-response because it is based on the scoring of all members of a sample population for having or not having a certain condition at a given applied dose (Casarett et al., 2008). This is precisely what is done through flow cytometry assessing a sample on a cell
by cell basis for being dead or alive. Using flow cytometry we can measure the fraction of cells killed \((fa)\) at a dose \(D\) with high accuracy and precision due to the large number of cells analyzed which ensures an extremely low standard error (SE). However, high accuracy and precision apply to a single sample and not to replicates. The source of variability between replicates will be both biological and experimental. For example cells overgrown in culture may respond with higher variability than cells in exponential growth when estimated from replicates. Similarly any problems around drug exposure or the staining procedure will add to the variability of replicates although the precision and accuracy of each sample determination will be very high due to the large amount of cells scored in each sample tube. Regarding calculation of the median cytotoxic dose \(D_m\) this replicates will have a critical impact on the statistical precision of the \(D_m\) estimate.

### 3.1 Calculating the median dose: The median effect equation

Cells cultured in vitro can be exposed to increasing doses of a cytotoxic drug during a certain time interval (e.g., 48h or 72h) to determine a median cytotoxic dose \(D_m\). Several doses should be tested to extend over a dose range. The lower doses should induce a fraction of dead cells close to that of unexposed cells, while the higher doses should induce death values approaching 100% or achieve a plateau of maximal effect. In between these boundary doses the more intervals assessed the more precision we will get in the estimates. For example seven doses assessed in triplicate that yield cell death fractions between 5% and 95% would be enough to obtain regression estimates with an adequate precision. The dose response sigmoidal curve can be fit to a two parameter logistic function of the type:

\[
fa = 1 / [1+1/(D/Dm)^m]
\]  

where \(D\) is the dose, \(D_m\) is the dose required to achieve the median cytotoxic effect, \(fa\) is the fraction of dead cells, and \(m\) is a measurement of the sigmoidicity of the curve. When \(m=1\) the dose-effect curve is hyperbolic, when \(m>1\) the curve is sigmoidal, while \(m<1\) indicates a negative sigmoidal shape (Fig. 3A).

To estimate \(D_m\) and \(m\) the median effect equation is written as:

\[
fa/(1-fa) = (D/Dm)^m
\]

The factor \((1-fa)\) is the fraction of live cells. Applying a log transformation the following linear function is obtained:

\[
\log (fa/(1-fa)) = m \cdot \log (D) - m \cdot \log (Dm)
\]

Thus plotting log values of experimental doses against log values of the ratio of dead/live cells will show a linear trend that is often referred to as median effect plot (Fig. 3B).

This is a linear function of the type \(y = mx + b\), where \(y = fa/(1-fa)\), \(x = \log(D)\), and \(b = -m \cdot \log (Dm)\). A linear regression can be applied to these data to obtain estimates for \(m\) and \(Dm\). The \(m\) coefficient can be readily determined by the slope of the regression, and \(Dm\) is derived from the estimate of the intercept \(-m \cdot \log (Dm)\).

The squared correlation coefficient or \(R^2\) value is an estimate of the precision of the overall regression (Fig. 3B). In this particular application to data representing cell death vs. dose of cytotoxic drug, \(R^2\) is affected mainly by the scattering of replicate values, which in turn
depend on the experimental and biological variability, and also on the number of sample doses in between the lowest and highest effect values. The standard error (SE) of \( m \) and the intercept can also be obtained from regression analysis to get a 95\% confidence interval around \( \log(Dm) \). The formula for manual calculation is rather complicated and involves computing \( SE(\log(D)) \) when \( D=Dm \) (shown in eq. 14, Table 5) but may be obtained using any software that implements this calculation such as CalcuSyn or Compusyn (Bijnsdorp et al., 2011; Ikeda et al., 2011; Ramachandran et al., 2010).

\[
y = 4.9392x - 4.1445
\]
\[
R^2 = 0.9014
\]

Fig. 3. Median effect plot. A. The two parameter dose-response sigmoidal curve for a particular example with \( Dm=7\mu M \) and \( m=5 \). B. Algebraic and log transformation to obtain a linear function. A linear regression can be applied to experimental data in order to obtain estimates of the two parameters \( Dm \) and \( m \). The squared correlation coefficient \( R^2 \) is a measure of the overall precision of the linear regression and thus the \( Dm \) and \( m \) estimates.

3.2 Threshold, median dose and maximal efficacy

Analysis of the dose-effect plot can be informative about threshold values and maximal efficacy values.

In practice the threshold value will be the minimal dose where the fraction of dead cells is higher than that of untreated cells. The maximal efficacy would be the fraction of dead cells where the sigmoidal curve approaches a plateau. Quite often the maximal efficacy approaches 100\%. However a cell population may exhibit a differential response and a fraction of cells may require quite larger doses. In these cases the maximal efficacy will be much less than 100\%.

3.3 Comparing two drugs

To compare two drugs and evaluate whether combination results in synergy or not a first step is to calculate \( Dm \) for the two drugs. Thus, the same approach described above should be applied to the second drug. The procedure will include evaluating several doses in replicates with the same exposure time interval as the first drug, obtaining data to create a median effect plot and estimating \( m \) and \( Dm \) by linear regression (Chang et al., 1987; Chou & Talalay, 1984; Sugiyama et al., 1998).
Results from assessment of the two drugs can be analyzed together in a combined median effect plot where log doses (in molar units) are depicted against log (fa/1-fa). In this plot the relative potency of the two drugs can be easily appreciated (Fig. 4A). A drug a is said to be more potent than a second drug b regarding the cytotoxic effect when less dose of drug a is needed to achieve the same cytotoxic effect with drug b on a molar basis. In addition when the slopes of the two drugs are different it suggests that the drugs have different mechanisms to induce cytotoxicity.

The dose effect equation can be re-written to calculate the dose required to induce a given cytotoxic effect:

\[ D = D_m \left( \frac{fa}{(1-fa)} \right)^{1/m} \]

For example the effective cytotoxic dose 50% (EC50) is the dose D that is estimated to kill 50% of the cells. In this case fa=0.5 and EC50 is coincident with the median dose Dm. Similarly EC25 is the dose D that is estimated to kill 25% of the cells.

### 3.3.1 Assessing the combined effect of two drugs under fixed molar ratio

Once obtained the dose-response curve for two drugs a and b, a third experiment with combination of a + b will be needed to determine if the interaction of these drugs is additive, synergic or antagonistic.

Assuming that the drug b is less potent than the drug a, fixed molar ratio could be used in the combination based on the relative potency EC50(a)/EC50(b). For example if EC50(a) is 10\( \mu \)M and EC50(b) is 30\( \mu \)M, the molar ratio of the combination would be 1/3. An empirical approach is to start the combination experiment with a combination of a+b calculated as

\[ EC50(a+b) = 10 \left( \frac{\log(EC50(a)) + \log(EC50(b))}{2} \right) \]

In this example this estimated value would be 17.3 \( \mu \)M. Assuming the fixed molar ratio 1/3 this combination would have 4.3\( \mu \)M of drug a and 13.0 \( \mu \)M of drug b.

Next we should treat this combination as a new drug a+b and evaluate several doses above and below 17.3\( \mu \)M in replicates to span a dose range of the combination. Thus, we will experimentally obtain a new data set of doses and cytotoxic effects that we should evaluate by the same procedure with the median effect plot, and conduct a linear regression to obtain estimates for m and Dm with the combination of a+b.

In particular we will obtain an effect-dose equation as shown above (eq. 4) to determine the dose of the combination a+b to achieve a desired cytotoxic effect level (EC)

\[ D = D_{m_{a+b}} \left( \frac{fa}{(1-fa)} \right)^{1/m_{a+b}} \]

For example applying (eq.5) EC50 (a+b) will be equal to the median dose estimated from the regression in the combined experiment (Dm_{a+b})

### 3.4 Graphical analysis

A first approach is to plot this result together with results of single drug effects to depict some relevant information (Pegram et al., 1999). When the combined-drug curve lies in a
midpoint between the two single drug curves it suggests an additive effect (Fig. 4A). It indicates that the potency of the combined drugs is at an intermediate point between the potency of each drug. When the combined drug curve is shifted to the left it would be closer to the more potent single drug and thus suggests synergism (Fig. 1B). On the other side when the combined drug curve is shifted to the right and closer to the low potency drug it is suggestive of antagonism (Fig. 1C). Another hint that could aid in generating hypothesis is the curve shape and particularly the slope. The variability around Dm is represented by the standard deviation (SD) of the Gaussian distribution underlying the quantal dose-response model discussed above. A large SD is in accordance with a flat curve while a small SD is in accordance with a steep curve (Fig. 2C,D). This variability has a biological significance and two drugs having different mechanisms of inducing cell death in a certain cell line may have different slopes.

### 3.5 Calculating the combination index (CI)

The graphical analysis gives some clues about what kind of interaction results from the combination of drugs a and b and depicts useful information but is less conclusive in quantitative terms. A more thorough conclusion can be derived from computing the combination index for each cytotoxic effect level (Chou, 2008; 2010). Computing the combination index (CI) for each effect level provides an answer to what kind of interaction occurs between drug a and drug b throughout the whole dose range.

The combination index method takes data provided by single and combined dose-effect equations to provide an estimate at the whole range of cytotoxic effects. The combination index is defined for a given effect level i by the following equation:

\[
CI (i) = \frac{Dac(i)}{Das(i)} + \frac{Dbc(i)}{Dbs(i)} + \alpha \frac{Dac(i) \cdot Dbc(i)}{Das(i) \cdot Dbs(i)}
\]

Where \(Dac(i)\) and \(Dbc(i)\) are the doses of drugs a and b respectively required in the combination a+b to produce an effect level i.

\(Das(i)\) and \(Dbs(i)\) are the doses of drug a and b respectively, required to produce an effect level i when used as single drugs. For any level i, these values are obtained from the three dose response curves defined by (eq. 4) (two single and one combined) obtained with parameters Dm and m that in turn were obtained from regression analysis with (eq.3) applied to experimental data. It is often assumed the conservative criteria that cytotoxic drugs are mutually non exclusive and \(\alpha=1\). If the three lines are strictly parallel and both drugs have a similar molecular target it could be assumed that they are mutually exclusive and in that case \(\alpha=0\). If the fixed molar ratio of drug a and b in the combined treatment is \(p/q\), then for an effect level i:

\[
Dac(i)=p/(p+q) \cdot Dm_{a+b} \cdot (fa(i)/(1-fa(i)))^{1/ma}
\]

\[
Dbs(i)=q/(p+q) \cdot Dm_{a+b} \cdot (fa(i)/(1-fa(i)))^{1/mb}
\]

\[
Das(i)=Dm_{a} \cdot (fa(i)/(1-fa(i)))^{1/ma}
\]

\[
Dbc(i)=Dm_{b} \cdot (fa(i)/(1-fa(i)))^{1/mb}
\]

\[
\alpha=\frac{Dac(i) \cdot Dbc(i)}{Das(i) \cdot Dbs(i)}
\]
Where \( fa(i) \) is the fraction of dead cells at effect level \( i \), \( D_{m_a} \) and \( m_a \) are the median dose and the slope estimated for drug \( a \), \( D_{m_b} \) and \( m_b \) are the median dose and the slope estimated for drug \( b \), and \( D_{m_{a+b}} \) and \( m_{a+b} \) are the median dose and the slope estimated for the combined treatment with drugs \( a \) and \( b \). Thus the combination index is calculated for any effect level above 0 and below 1.

\[
y = 3.2019x - 4.7512 \\
R^2 = 0.9272
\]

\[
y = 3.3544x - 3.4487 \\
R^2 = 0.8798
\]

\[
y = 3.0455x - 3.7415 \\
R^2 = 0.9852
\]

Fig. 4. Drug interaction and median effect plot. A. In this example experimental values are represented by circles and linear regression is applied to obtain estimates for \( D_{m_a} \) and \( m_a \). Experimental values obtained for a drug \( a \) are shown in black circles. The values of \( D_{m_a} \) and \( m_a \) for drug \( a \) were 10 \( \mu \)M and 3.0 respectively. Experimental values for a less potent drug \( b \) are shown in open circles. The values of \( D_{m_b} \) and \( m_b \) for drug \( a \) were 30 \( \mu \)M and 3.0 respectively. A combined experiment was run with \( a+b \) with a constant mass ratio of 1/3 based on \( EC_{50}(a)/EC_{50}(b) \) and assuming that under additive effect \( EC_{50}(a+b) = 10\left(\log(EC_{50}(a)) + \log(EC_{50}(b))\right)/2 \). Experimental values for the combination are shown in grey circles. The values of \( D_{m_a} \) and \( m_{a+b} \) for the combination \( a+b \) were 17.0 \( \mu \)M and 3.0 respectively. In this example drugs \( a \) and \( b \) have an additive interaction and the median effect plot of the combination lies in a mid position between the lines corresponding to the single drugs. B. The same experiment shown in A, but in this case the drugs \( a \) and \( b \) have synergic effect. The values of \( D_{m_a} \) and \( m_{a+b} \) for the combination \( a+b \) were 14.0 \( \mu \)M and 3.0 respectively. The median effect plot of the combination is shifted towards the drug \( a \), which has the highest potency. C. The same experiment shown in A, but in this example the drugs \( a \) and \( b \) have antagonistic effect. The values of \( D_{m_a} \) and \( m_{a+b} \) for the combination \( a+b \) were 24.4 \( \mu \)M and 2.9 respectively. The median effect plot of the combination in this case is shifted towards the drug \( b \) which has the lowest potency.
### Table 5. CI calculation between two drugs a and b at the 50% effect level under three alternative conditions: additive, synergic, antagonistic. To obtain CI as a function of the effect level $i$, the calculation has to be repeated for each arbitrary level $i$ between 0 and 1. A 95% confidence interval around $D$ in general and around $D_m$ in particular, can be computed using the formulas for standard error (SE, eq. 13 and eq. 14). A 95% confidence interval around CI at any effect level $i$ can be computed from the standard error formulas presented in eq. 12, 13 and 14.

| Single drug a | Single drug b | Combined drugs $a+b$ | Combined drugs $a+b$ | Combined drugs $a+b$ |
|---------------|---------------|----------------------|----------------------|----------------------|
| $D_m$ (uM) $(t)$ | 10            | 30                   | 17                   | 14                   | 24.4                 |
| $m$ $(t)$     | 3             | 3                    | 3                    | 3                    | 3                    |
| $f_{ai}$      | 0.5           |                      |                      |                      |                      |
| $D_{as}$ $(t)$| 10            |                      |                      |                      |                      |
| $D_{bs}$ $(t)$| 30            |                      |                      |                      |                      |
| $D_{ac}$ $(t)$| 12.75         |                      |                      |                      |                      |
| $p/(p+q)$ $(t)$| 0.25          |                      |                      |                      |                      |
| $q/(p+q)$ $(t)$| 0.75          |                      |                      |                      |                      |
| CI $(i)$ $(t)$ | 1.03          |                      |                      |                      |                      |

SE(CI$(i)) = \left( \frac{D_{ac}(i)/D_{as}(i) + D_{bc}(i)/D_{bs}(i)}{(SE(D_{ac}(i)/D_{ac}(i)) + SE(D_{as}(i)/D_{as}(i)))^2} \right)^{1/2}$

(eq. 12)

SE($D$) $= 1/2 \cdot 10^{[\log(D) + SE(\log(D))]} - 10^{[\log(D) - SE(\log(D))]}$ (eq. 13)

SE($\log(D)$) $= (\log(D) + \frac{SE(b)}{\log(f_{ai} / (1-f_{ai}) - b)} + \frac{SE(m)}{m} + 2[(\log(D))^2 - 2 \cdot (\log(D))] / (\log(D) / (\log(b) - \log(m)m/h)^{1/2}$

(eq. 14)

where $b = m \cdot \log(Dm)$

# (1) Obtained from linear regression of experimental data

# (2) $D_{as}(i) = D_{m1} \left( \frac{f_{ai}(i)}{1-f_{ai}(i)} \right)^{m_{1a}}$

# (3) $D_{bs}(i) = D_{m2} \left( \frac{f_{ai}(i)}{1-f_{ai}(i)} \right)^{m_{2b}}$

# (4) $D_{ac}(i) = p/(p+q) \ D_{m1} \left( \frac{f_{ai}(i)}{1-f_{ai}(i)} \right)^{m_{1a}} + b$

# (5) $D_{bc}(i) = q/(p+q) \ D_{m2} \left( \frac{f_{ai}(i)}{1-f_{ai}(i)} \right)^{m_{2b}} + b$

# (6) Molar ratio $p/q = D_{m1} / D_{m2} = 10/30 = 1/3$

# (7) CI$(i) = D_{ac}(i) / D_{bs}(i) + D_{bc}(i) / D_{bs}(i) + \alpha D_{ac}(i) . D_{bc}(i) / D_{as}(i) . D_{bc}(i)$

A 95% confidence interval around $D$ in general and around $D_m$ in particular, can be computed using the formulas for standard error (SE, eq. 13 and eq. 14). A 95% confidence interval around CI at any effect level $i$ can be computed from the standard error formulas presented in eq. 12, 13 and 14.
Table 5 summarizes a manual calculation of CI of two drugs a and b using these formulas for the 50% effect level under three hypothetical results: additive, synergic or antagonistic effect.

The same calculation can be applied to any effect level to plot CI as a function of effect level. When the interaction is additive CI = 1. In this case it can be interpreted that one of the drugs (the less potent one, i.e. drug b in the example) is acting as though it is merely a diluted form of the other (drug a in the example). When CI < 1 the combination of a+b is synergic while CI > 1 indicates antagonism. Synergy implies that the combination of the two drugs achieves a cytotoxic effect greater than that expected by the simple addition of the effects of the drugs a and b, while antagonism achieves a cytotoxic effect lower than that expected by additive effects of drugs a and b.

Fig. 5. Drug interaction and CI calculation. A. Only rarely the combination index obtained is constant for all effect levels. Here it is shown how different values of the slope m obtained through linear regression in the combination experiment (a+b) would affect the shape of the curve representing the CI as a function of the effect level. Similarly, differences between the slopes obtained for drugs a and b through the single drug experiments will contribute to the uneven shape of the CI function. Note that depending on the effect level the interaction a+b with m=1.8 would be synergism, additive or antagonism at EC25, EC50, and EC75 respectively (arrows). B. Results of CI calculation for the example where a+b results in synergism considering m=4 in the regression of the combined-drug experiment. C. Results of CI calculation for the example where a+b results in antagonism considering m=2.5 in the regression of the combined-drug experiment. D. 95% confidence level intervals around CI, using an algebraic approximation (eq. 12, Table 5) in an example where combination of a+b is synergic.
The horizontal line corresponding to $CI(i)=1$, where $i$ is any effect level in the interval $(0,1)$, is often call the additive effect line. A combination of drugs $a$ and $b$ may result in CI values above or below the additive line at different effect levels. Thus, the CI as a function of effect level is not constant or linear and it may be decreasing or increasing (Fig 5A). If data from the combination experiment in the example of figure 4A resulted in $m_{a+b}=5$ or $m_{a+b}=1.8$, even still with $Dm_{a+b} = 17$ the CI line would be inclined downwards or upwards respectively (Fig. 5A). Only at effect levels close to EC50 the result would be strictly additive. An important conclusion is that for some drug combinations, experiments conducted at different single dose-effect levels may yield opposing results. For example if the combination $a+b$ with $m=1.8$ shown in figure 5A were experimentally evaluated only at effect level 0.25 the single dose analysis would conclude on synergism. However if it were evaluated at effect level 0.75 it would conclude on antagonism (Fig. 5A). This exemplifies why the assessment of combination index over the whole dose range will show all kinds of interactions that may result from combination at different effect levels.

Computing a standard error of CI allows plotting confidence intervals at all effect levels providing a further assurance over the computation. A 95% confidence interval will indicate that if we repeat the experiment 100 times, 95 out 100 times the CI would be within this interval. For example, observing whether or not confidence limits are above or below the additive line will allow concluding with further statistical support on antagonism or synergism respectively. Computation of the standard error of CI and confidence intervals at all levels should be better obtained through specialized software such as Calcusyn or Compusyn (Bijnsdorp et al., 2011; Chou, 2010). It may also require approaches such as Monte Carlo simulation based on the estimated parameter for $m$ and $Dm$ in single and combined equations.

4. Concluding remarks

A thorough assessment of drug interaction is an essential step in targeted combined therapy. The new targeted agents are seldom useful as single agents but may be effective when used in specific combinations. The median effect and combination index calculation are well founded methods traditionally used in pharmacological and toxicological studies. Since new cytotoxic drugs target mechanisms eliciting cell death, biomarkers related to viability assessment are preferred to biomarkers of cell proliferation. Flow cytometry is an ideal technology to provide massive data from cell death biomarkers to build dose response curves of cytotoxic effect. When these data is further used to determine the combination index a full characterization of drug interaction over the cytotoxic effect is obtained at all effect levels. This approach can be applied to tumor cell lines in preclinical studies and also in patient-derived tumor cells, thus providing useful information as prospective indicators of the potential therapeutic response to combined-drug antitumor treatment.

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