Photon Emissions as Differential Indicators for Different Components of Protein Kinase A (PKA) in Transfected Murine Melanoma Cells

Blake T Dotta, Carly A Buckner, Robert M Lafrenie and Michael A Persinger

Biomolecular Sciences Program, Laurentian University, Sudbury, Ontario, Canada P3E 2C6

Corresponding author: Michael A Persinger, Biomolecular Sciences Program, Laurentian University, Sudbury, Ontario, Canada, Tel: 01-705-675-4824; Fax: 01-705-671-3841; E-mail: mpersinger@laurentian.ca

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Abstract

Increased emissions of photons or shifts in spectral power densities of photons have been reliably measured from malignant cells compared to non-malignant cells. Previous experiments have shown that specific wavelengths within the visible spectrum emitted from melanoma cells were associated with the activation or inhibition of specific molecular structures or pathways. To discern if numbers of photons could differentiate the dynamic state of a critical protein (enzyme), Protein Kinase A (PKA), melanoma cells were transfected with either catalytic subunits, regulatory subunits, or a mutant dominant negative PKA. Compared to typical melanoma cells those transfected with the regulatory subunit exhibited a marked (10 to 100) increase in photon emissions for several hours. The small but significant increase in photon emissions from cells transfected with the catalytic subunit was more brief (first 20 min) and less intense. Photon emissions from cells transfected with inhibitory components did not differ from typical melanoma cells. The vectorial characteristics of the photon emissions were sufficient to clearly differentiate activation of various components of PKA domains.

Keywords: Ultraweak photon emissions; PKA; Regulatory domain; Catalytic domain; Inhibitor domain; Melanoma cells; Intercellular communication

Introduction

There are a variety of signaling pathways involved with the induction of conditions that promote aberrant cellular division and tumor formation. Although the numbers of different types of cells that can be differentiated by histological criteria are within the hundreds, the myriad of experiments during the last half-century of molecular biology have suggested common pathways [1]. The identification of these pathways is often tedious, labour-intensive and expensive. Several recent experiments [2-4] have demonstrated that the measurement of photon emissions from malignant cells and the extraction of the spectral power densities of the amplitude-modulation of the visible band can differentiate normal and abnormal conditions. Quick measurements of photon profiles that discern components of activation from regulatory subunits of primary molecules in signaling pathways may allow alternative methods to monitor cellular states. Here we present clear evidence that transfection with different subunits of Protein Kinase A (PKA) which differentially regulates PKA activity and impact major pathways involved with cellular function produced differential magnitude and temporal patterns of ultraweak photon emissions from transfected melanoma cells.

Photons exhibit unique features that constitute properties of both waves and particles. Theoretically photons have the capacity to mediate the interface between matter whose functions are determined by spatial (molecular) structure and energy whose functions are determined by the temporal pattern of the sequences [5]. Dotta et al. [3], who employed Cosic’s model involved with recognition by molecular resonance, demonstrated a clear relationship between specific wavelengths of photon emissions from mouse melanoma cells and specific molecules that either inhibited or facilitated these emissions. During the early 20th century Gurwitch [6] had postulated that all living systems emit photons. Popp [7] extended this concept and suggested that since all energy that presently composed living systems originated from the sun, “virtual” photons might be stored within every cell. Persinger [8] showed quantitatively that the ubiquitous approximately $10^{-12}$ W·m$^{-2}$ of photon flux density emitted by living systems was consistent with a fraction of spontaneous ‘re-emission’ of the accumulation of solar photon energy since Life evolved. Trushin [9] Fels [10] and Bajpai et al. [11] have argued that these photon emissions from cells and bacteria may reflect the fundamental method of information transfer. Like action potentials whose energy is contained within the axonal membrane and not carried by the propagating spike, the photons between cells contain the information or “keys” that initiate the powerful forces and energies that are self-contained with the molecular pathways.

Protein Kinase-A [1] has been described as a family of enzymes whose primary activities are dependent upon the concentrations of cyclic AMP within cells. Cell-modulating molecular structures circulating within the vasculature such as adrenalin and glucagon initiate the cascade that triggers the dynamics of PKA. Four cyclic AMP molecules activate a single
PKA macromolecule that augments but does not participate directly in subsequent reactions. A pair of cyclic AMP molecules is sequestered by each of two regulatory subunits which detach from the catalytic subunits that become activated. The catalytic subunits ultimately contribute to the phosphorylation of other proteins. From some perspectives PKA markedly influences most if not all of the cellular responses associated with the cyclic AMP as well as other specific enzymes, ion channels (particularly calcium channels), H1 (histones) and CREB (cyclic AMP response element binding factor) involved with promoter selections that stimulate transcription. Considering the role PKA plays in malignant and non-malignant cells and the availability of different subunits, this enzyme was considered optimal to explore photon emission correlates.

**Method**

All experiments and each treatment involved a minimum of three replicates and were conducted over a period of several months. B16-BL6 murine melanoma cells [12,13] were transiently transfected with PKA-CA (MGC-6169, Clontech), PKA-REG (MGC-11864, Clontech), or PKA-DN (Addgene 16716). Background readings were obtained from the vector only control mCherry vector (Clontech) condition [14]. Only one transfected condition was tested per day. The PKA-DN constructs corresponded to the regulatory domain that contained the mutations in the cAMP binding sites.

The experimentally manipulated or control melanoma cells were maintained on cell culture plates as specified [15]. The cells were grown to approximately 80-90% confluence 48hr post-transfection when the photon readings were measured (corresponding to approximately 10^6 cells per dish). Due to equipment availability photon measurements were obtained for only one plate per day. The PKA-DN constructs corresponded to the regulatory domain that contained the mutations in the cAMP binding sites.

The average photon counts per second for the first hour of measurement for the control (typical melanoma), PKA-Dominant Negative (DN), PKA Catalytic Subunit (CA) and PKA Regulatory Subunit (REG) expressing plasmids are shown in Figure 3. The values for the mCherry empty plasmid control (not shown) produced no significant differences from baseline values.

**Results**

For part 1 of the study that involved the measurement of photon emissions for 22.5 hrs after removal from the incubator, the results were conspicuous as shown in Figure 1. Those cells transfected with regulatory or catalytic domains of PKA units exhibited markedly increased photon counts for the first approximately 300 s (5 min). On the other hand the typical melanoma cells (untransfected) and control vector transfected melanoma cells displayed <50 counts per s. More specifically during the first 5 min there was a factor of 200 increase in photon emissions from the PKA transfected cells compared to the control cells with PKA activity.

![Figure 1](image1.png) **Figure 1** Numbers of photons per second per 2.5 increments for PKA transfected melanoma cells (grey) compared to typical melanoma cells (black line). Horizontal axis indicates serial order of 2.5 s increments.

On the other hand the cells that were transfected with the dominant negative inhibitory PKA plasmid, a type of control for experimental manipulation of cell dynamics, did not display the massive increase in photon emissions. As shown in Figure 2, the photon emissions ever over 22.5 hrs was within the typical range of melanoma cells removed from incubation. The only discrepancy was the burst that occurred about 4 hrs later. However the magnitude was not comparable to the photon emissions measured during the first 5min from the cells transfected with the regulatory or catalytic PKA domains.

![Figure 2](image2.png) **Figure 2** Example of numbers of photon counts per s for a continuous series of 2.5s increments (horizontal axis) from melanoma cells transfected with the Dominant Negative components.

The average photon counts per second for the first hour of measurement for the control (typical melanoma), PKA-Dominant Negative (DN), PKA Catalytic Subunit (CA) and PKA Regulatory Subunit (REG) expressing plasmids are shown in Figure 3. The values for the mCherry empty plasmid control (not shown) produced no significant differences from baseline values.
conditions (cells with no treatment). One way analyses of variance demonstrated a statistically significant difference \[ F(3,22)=3.20, p<0.05; \text{eta}^2=0.33 \] in numbers of photon emissions between the groups. Stated alternatively one-third of the variance in photon emissions were explained by the experimental manipulations.

The more protracted display of photon emissions over 9, 20 min blocks as a function of the different experimental conditions as well as the effects of the chemotherapeutic agent are shown in Figure 4. Again the marked increase in average photon counts per 20 min block for the first 40 min was emitted by the enhanced regulatory unit transfection compared to all other groups. The effects of the drug increased photon counts compared to all other treatments for the first 40 min, including the negative control (DN). Clearly the most significant enhancement of photon emissions occurred for the cells transfected with the regulatory subunit which was evident for at least 3 hr.

To demonstrate protein expression after 48 hours, a Western blot analysis was conducted and can be seen in Figure 5. This clearly illustrates the lower exposure time of Vector-only and PKA-CA when compared to PKA-DN and PKA-REG.
Discussion

If photons contain the information that control the vectorial characteristics of inter-cellular or intraorganismic interactions, then the numbers of photons should be qualitatively and quantitatively correlated with those molecular components that are known to control the dynamics of a cell. We have assumed that the actual massive amounts of energy that control the molecular machinery are contained with the glucose metabolism-based chemical processes that are determined by the genetics of the cell. The photons would be the local and non-local determinants of the information that initiates or terminates these processes. One metaphor is that the energy and forces required to move a significant mass such as an automobile are substantial. However its initiation requires the small effort of turning the key in the ignition and its direction involves a small effort to turn the steering wheel.

Previously Murugun et al. [12] showed that injecting morphine into typical melanoma cells produced brief bursts of photons. The magnitude was considered sufficient to facilitate the known metastases associated with this opiate. Karbowski et al. [13] showed that only the introduction of melanoma cells into normal cells such that the former constituted only 1% of the population was sufficient to increase photon emissions. This flux density would be sufficient to affect substantial distances particularly if non-locality was involved [16]. Dotta et al. [3] had shown that during states as the melanoma cells adjusted to ambient temperatures there were predictable shifts in the peak wavelength of the photon emissions across the visible spectrum. All of these experiments support the hypothesis that photons may be the source of information that triggers the determinant pathways that lead to malignancy and its proliferation.

In the present study melanoma cells transfected with PKA demonstrated an extraordinary enhancement of photon emissions as shown in Figure 1. Assuming the average of 8000 photons per s during the first 6 to 8 min, this would be equivalent to more than $10^{-11} \text{W} \cdot \text{m}^{-2}$. This is at least a factor of 10 greater than photon flux densities emitted by typical melanoma cells. On the other hand cells transfected with the PKA dominant negative vectors where PKA was inhibited by the transfection, showed photon emissions in the order of 300 photons per s or about $10^{-12} \text{u} / \text{m}^2$, which is more typical of ultraweak photon emissions from typical melanoma cells and even from non-malignant brain slices [17].

The implications of the differential photon emissions between the regulatory and catalytic components of PKA are significant. The regulatory subunit normally interacts with the catalytic domain of PKA to inhibit activity but when it sequesters cyclic AMP it dissociates from the catalytic domain that then becomes capable of enzymatic activity. There is also strong evidence that the regulatory subunits can exhibit additional biological activity. Considering these functions the markedly increased photon emissions when cells were transfected with this component as well as the protracted emission would be expected. The catalytic subunit, the active site for PKA, which contains a domain to sequester the molecules for phosphorylation (and binds ATP) would be expected to exhibit more brief and discrete activity which would require less photon density. This inference was supported by the measurements. In summary these results indicate that the temporal profile of photon flux density might be employed as a tool to discern which component of a signaling pathway has been differentially activated.

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