# **STUDIA UNIVERSITATIS** BABEŞ-BOLYAI



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2

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#### SUMAR - CONTENTS - SOMMAIRE - INHALT

S. TOT, S. OROIAN, Prediction of Vegetation Dynamics on the Tailing Dumps in Căpuş and Aghireş (Cluj County) Based on the Researches Performed During the 2005 – 2011 Period	5
B. DRUGĂ, V. BERCEA, C. SICORA, The Effect of Various Light Intensities on Photosystem II Photochemistry in the Cyanobacterium <i>Synechocystis</i> sp. PCC 6803	15
A. HEGEDŰS, V. BERCEA, C. SICORA, The Study of Photosystem II (PS II) Activity Based on Chlorophyll Fluorescence in <i>Aphanizomenon elenkinii</i> AICB 709, under Light Stress Conditions.	35
Z. TANOURI, O. RABIEIMOTLAGH, An Estimation for Lower Bound of p53 in DNA Healing Process; A Mathematical Approach	. 53
IN MEMORIAM: Centenarul nașterii Profesorului George Emil Palade	71
BOOK REVIEW: Ecological and Behavioral Methods for the Study of Bats, 2 <sup>nd</sup> Edition.	. 81

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#### Original picture on front cover:

Microscopic view of the filamentous cyanobacterium Anabaena strain ATCC29413

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# PREDICTION OF VEGETATION DYNAMICS ON THE TAILING DUMPS IN CĂPUȘ AND AGHIREȘ (CLUJ COUNTY) BASED ON THE RESEARCHES PERFORMED DURING THE 2005 – 2011 PERIOD

# SABINA TOT<sup>1</sup> and SILVIA OROIAN<sup>2</sup> $\boxtimes$

SUMMARY: The paper presents the floristic structure of the nodal phytocenoses identified in the vegetation dynamics on the tailing dumps from Căpuş and Aghireş (Cluj County). The incipient vegetation phase of the tailing dumps, regardless of the chemical nature of the stored tailings, is represented by the Tussilago farfara phytocenoses. Within the following phase, when certain Poacee species begin to grown on the tailings, the phytocenoses succession is different. On the acid tailing dumps, one may notice the *Calamagrostis* epigejos and the Festuca rubra phytocenoses with Agrostis capillaris, while on the neutral or alkaline reaction tailing dumps the Elymus repens and Festuca rupicola phytocenoses are installed. These phytocenoses keep their individuality until the populations of some arbuscular species present in their structure, become prevalent. Within the current study, the Crataegus monogyna and Prunus spinosa phytocenoses end the succession of herbal phytocenoses and enable the growth of forest phytocenoses by the created microclimate.

Keywords: tailing dumps, phytocenoses, vegetation succession.

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#### Introduction

The two tailing dumps were formed due to the economic activities developed in this region for the valorization of mineral substances used in the industry. The tailing dumps from Căpuş developed as a result of tailing deposition during the 1969 – 1975 period, pursuant to the processing of the iron ore mined from the North-Eastern side of the Gilău Mountains. The tailing dumps from Aghireş were formed as a result of kaolin exploitation in the Northern part of the Dumbrava locality (Pop, 2001, 2007). The tailing dumps in Căpuş, located at 540-590 m.s.m. are 20 - 25 years old and they have a clay-sandy-rocky structure, forming an acid reaction technogenous soil, while the tailing dumps in Aghireş, located at of 450-480 m elevation, are 20-25 years old, with a sandy-clayey structure, and a weak-alkaline or neutral reaction.

The vegetal communities developed on the two types of tailing dumps have followed a partially different dynamics.

#### Materials and methods

The research of the vegetation dynamics on the tailing dumps employed the inventory squares method (Anghel *et al.*, 1971, Cristea *et al.*, 2004) and the method of the phytocenological surveys.

In order to grasp the evolution of the vegetation dynamics, the authors have performed quantitative field measurements, by means of the metric frame, in 120 inventory squares and have performed 292 phytocenological surveys on the tailing dumps in Căpuş and Aghireş, under various vegetation stages.

#### VEGETATION DYNAMICS ON THE TAILING DUMPS)



Figure 1. Tussilaginetum farfarae on Western slope of sampled site.

#### **Results and discussions**

The floristic composition of phytocenoses installed on the tailing dumps differ depending on the nature of the exploited geologic substrata and on the duration of the deposits formation.

Within the first stage of plant colonization, regardless of the geological structure, the *Tussilago farfara* phytocenoses accommodate. Their structure includes the following species: *Leontodon autumnalis, Convolvulus arvensis, Cirsium arvense, Setaria viridis, Trifolium repens, Polygonum persicaria, Poa compressa, Sonchus arvensis* (Table 1, column 1).

The second stage, consisting in tailing colonization, begins with the occurrence of some species of Poacee perene. These are installed on the tailings sections where some of the exceeding elements (Fe, Al, K<sub>2</sub>O, Mg, S) are leached. Starting with this stage, the dynamic evolution of the phytocenoses differs in relation to the physico-chemical and nutritional properties of the soil layer.

The Tussilago farfara pioneer communities developed on the tailing dumps in Căpus, whose reaction was mainly acid, due to the high content of SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub> and FeO (Despu, 1985), were followed mainly by the Calamagrostis epigejos oligotrophic phytocenoses. Together with this species, which covers 25-35% on average, highly present are the following species: Agrostis capillaris, Daucus carota ssp. carota, Plantago lanceolata, Cynosurus cristatus, Agrimonia eupatoria and Centaurium pulchellum (Table 1, column 2a). In a few years (5-6), when the tailing dump begins to turn into soil and when more nutrient substances accumulate at the bottom, representative substances develop, presenting some mesophile and mesothophic pastures phytocenoses. The prevailing plant species in these phytocenoses are: Phleum pratense, Arrhenatherum elatius and Dactylis glomerata, which cover together 35-45% (Table 1, column 4a). Beside these, the following species are frequently met: Agrostis capillaris, Daucus carota ssp. carota, Cynosurus cristatus, Achillea millefolium, Briza media, Centaurea nigrescens, Geranium pratense, Medicago falcata, Trifolium pratense, Trisetum flavescens, Leucanthemum vulgare, Lotus corniculatus and Centaurea jacea. These species reveal the mesophile and mesothophic nature of the pastures.

Towards the upper part of the tailing dump, where the nutritional elements are fewer, the *Agrostis capillaris* and *Festuca rubra* phytocenoses begin to develop. These phytocenoses expand more and more on the tailing dumps, until they cover their middle and upper part (Table 1, column 3a).

The economic non-use of these herbal phytocenoses will facilitate the development of *Prunus spinosa* and *Crataegus monogyna* arbuscular populations coming from the edge of the forest on the tailing dumps. The development of arbuscular phytocenoses on these tailings represents a final stage in the succession of the herbal vegetation and precedes the development of the forest (Table 1, column 5).

#### VEGETATION DYNAMICS ON THE TAILING DUMPS)

The *Tussilago farfara* pioneer communities developed on the tailing dumps in Aghireş, due to the clayey-sandy substrata, presenting an alkaline or neutral reaction, are followed first by *Elymus repens* xero-mezophile phytocenoses. The structure of these phytocenoses, together with some xerophile species such as: *Elymus hispidus, Euphorbia cyparissias, Centaurea micranthos, Agrimonia eupatoria, Fragaria viridis* includes also some mezophile species such as: *Achillea millefolium, Cynosurus cristatus, Holcus lanatus, Medicago lupulina, Leucanthemum vulgare* and *Prunella vulgaris* (Table 1, column 2b).

Location of the tailing dump	Căpuş (a)		Aghireş (b)					
The vegetation phase	1	2a	3a	4a	2b	3b	4b	5
Number of phytocenological	17	6	15	8	5	8	18	7
surveys								
Tussilago farfara	V	III	III			Ι	IV	III
Poa compressa	IV		Ι		II	Ι		
Calamagrostis epigejos	Ι	V	Ι		III		Ι	Ι
Agrostis capillaris	Ι	IV	V	V		II	V	Ι
Festuca rubra	Ι		V	Ι		V		Ι
Arrhenatherum elatius				V		Ι		III
Elymus repens	Ι		Ι		V		Ι	Ι
Convolvulus arvense					II			II
Galium verum		Ι	Ι	III	Ι	III	Ι	II
Festuca rupicola							V	
Crataegus monogyna	•	II			Ι		II	V
Prunus spinosa		Ι			Ι		Ι	V
Achillea millefolium	Ι	III	III	II	III	III	IV	
Achillea setacea		Ι	•	•	II	II		•
Agrimonia eupatoria	Ι	IV	III	III	III	II	IV	V
Agrostis stolonifera	II	IV	Ι		V	II	Ι	II

**Table 1.** The floristic structure of the nodal vegetal communities developed on the tailing dumps in Căpuş and Aghireş.

## Table 1 (continued)

Location of the tailing dump	Căpuş (a)		Aghireş (b)					
Anthoxanthum odoratum		Ι	Ι	Ι			Ι	V
Astragalus glycyphyllos				Ι			Ι	Ι
Brachypodium sylvaticum								III
Campanula patula			Ι	Ι			Ι	
Carex humilis							Ι	Ι
Centaurea jacea			Ι	III	II	Ι	Ι	II
Centaurea micranthos	II	Ι	Ι	III	III	III		II
Centaurea nigrescens			II			•		II
Centaurium pulchellum	Ι	IV			II	II	Ι	
Cichorium inthybus	Ι	III	Ι	Ι	II	II	Ι	
Cirsium arvense	Ι	II	Ι				III	Ι
Cerastium holosteoides			Ι	Ι		Ι	Ι	
Conyza canadensis	Ι	II	Ι		III	III		
The vegetation phase	1	2a	3a	4a	2b	3b	4b	5
Number of phytocenological	17	6	15	8	5	8	18	7
surveys								
Coronilla varia				Ι		Ι	Ι	II
Cornus sanguinea				Ι	Ι	Ι		Ι
Crepis biennis			Ι	Ι	II	Ι	Ι	
Cynodon dactylon			Ι			Ι		
Cynosurus cristatus	Ι	III	Ι	II	III	Ι	Ι	II
Dactylis glomerata		III	Ι	IV	•	Ι	Ι	V
Daucus carota ssp.carota	II	III	Ι	III	•	II	IV	
Dianthus carthusianorum				III	•	Ι	•	
Echium vulgare				III				
Elymus repens	Ι		Ι		V		Ι	Ι
Euphorbia cyparissias			Ι	Ι	II	Ι		
Fragaria viridis	Ι	IV	Ι	Ι	II	Ι	Ι	II
Galium mollugo	Ι		Ι	III	II	Ι	Ι	II
Geranium pratense				II				Ι
Hieracium pilosella	II	III	II		II	Ι	IV	
Holcus lanatus	Ι	III	Ι		IV	Ι	III	
Hypericum elegans	•		•	III				Ι

### Table 1 (continued)

Location of the tailing dump	Căpuş (a)		Aghireş (b)					
Hypericum perforatum		II	Ι	II	II	Ι	III	
Inula saalicina							Ι	
Leontodon autumnalis	Ι	IV	III	Ι	II	III	IV	
Leontodon hispidus			Ι	Ι		II	II	
Leucanthemum vulgare	Ι		Ι	II	II	II	Ι	
Lolium perene	•		Ι	Ι		Ι	Ι	
Lotus corniculatus	•		II	IV	II	II	IV	II
Medicago falcata	•		•	IV	Ι	Ι		II
Medicago lupulina			Ι	Ι	II	II	Ι	
Petrorhagia prolifera				II		Ι	Ι	
Phleum pretense				III	Ι	II		III
Plantago lanceolata	Ι	V	III	Ι	III	III	II	III
Plantago media	Ι	III		III		Ι	Ι	Ι
The vegetation phase	1	2a	3a	4a	2b	3b	4b	5
Number of phytocenological	17	6	15	8	5	8	18	7
surveys								
Poa pratensis			Ι	Ι		II	Ι	II
Polygonum persicaria	Ι	II	Ι		II	Ι		
Prunella vulgaris		III	Ι		II	II	II	III
Pyrus pyraster		Ι	II	Ι	Ι			
Ranunculus acris		Ι	Ι	Ι			Ι	IV
Ranunculus repens	Ι	III		Ι	III			Ι
Rosa canina		II		•	•		II	IV
Rubus caesius	•	III	•	•	•	•	III	Ι
Rumex acetosella	III		Ι	•	II		Ι	
Trifolium arvense				•	II	Ι	III	
Trifolium campestre	•			III	•	Ι	Ι	II
Trifolium pratense		Ι	II	III	III	Ι	IV	
Trifolium repens	II	III	II		II	Ι	IV	II
Taraxacum officinale	Ι	II	Ι	II		Ι	II	
Vicia cracca				III		•	Ι	II
Vulpia myuros	II	III	Ι		Ι	Ι	Ι	

#### S. TOT, S. OROIAN

Nodal phytocenoses in the vegetation succession on the tailing dumps

- 1. Tussilago farfara cenoses;
- 2a. Calamagrostis epigejos cenoses;
- 3a. Agrostis capillaris with Festuca rubra cenoses;
- 4a. Arrhenatherum elatius cenoses;
- 2b. Elymus repens cenoses;
- 3b. Festuca rubra cenoses with Galium verum;
- 4b. Festuca rupicola cenoses with Agrostis capillaris;
- 5. Prunus spinosa cenoses with Crataegus monogyna.

During the second succession stage, when the tailing dump begins to be structured as soil, the *Festuca rupicola* xerophile phytocenoses are mainly developed. The floristic composition of these phytocenoses includes a high prevalence of xerophile species among which: *Carex humilis, Coronilla varia, Astragalus glycyphyllos, Fragaria viridis, Galium verum, Hieracium pilosella, Inula germanica, I. salicina, Leontodon hispidus, Trifolium medium, Hypericum perforatum* and *Dianthus carthusianorum* (Table 1, column 4b). Together with these xerophile phytocenoses, in Aghireş there are also small areas of *Festuca rubra* mezophile phytocenoses with *Galium verum* and *Festuca rubra* with *Agrostis capillaris*. These are found mainly on the less steep slopes and on the small plateaus of the tailing dumps (Table 1, column 3b).

The floristic structure of these herbal phytocenoses rarely includes several arbuscular species, such as: *Crataegus monogyna, Rosa canina, Prunus spinosa* and *Ligustrum vulgare*. These will continuously develop and will form compact phytocenoses which will replace the herbal phytocenoses on the tailing dumps.

The final phase of the herbal phytocenoses is also represented in Aghireş by the *Prunus spinosa* and *Crataegus monogyna* phytocenoses (Table 1, column 5).

#### Conclusions

1. The vegetation of the tailing dumps is a long-term process and it is influenced by the physico-chemical properties of the tailings.

2. The first phytocenoses developed on the tailing dumps, regardless of the chemical nature of the substrata, are those developed by *Tussilago farfara*.

3. While the tailing dump begins to form as "technogenous soil" and it accommodates more and more Poacee species, the *Calamagrostis epigejos*, *Elymus repens*, *Festuca rubra* and *Festuca rupicola* mezophile and xero-mezophile phytocenoses begin to occur, which significantly contribute to the development of soil.

4. Finally, such herbal phytocenoses are replaced by *Crataegus monogyna* and *Prunus spinosa* arbuscular phytocenoses, which facilitate the development of forest phytocenoses.

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# THE EFFECT OF VARIOUS LIGHT INTENSITIES ON PHOTOSYSTEM II PHOTOCHEMISTRY IN THE CYANOBACTERIUM *SYNECHOCYSTIS* sp. PCC 6803

## BOGDAN DRUGĂ<sup>1</sup>, VICTOR BERCEA<sup>1</sup> and COSMIN SICORA<sup>1,2</sup>

SUMMARY: The cyanobacterium Synechocystis sp. PCC 6803 grown on BG 11 medium under air bubbling was exposed to high light intensities of 2000, 3000 and 4500 µmol.m<sup>-2</sup>.s<sup>-1</sup> while being in the exponential growth phase. The effect of light treatment was assessed after 15, 30, 45, 60, 75, 90, 105 and 120 minutes by monitoring chlorophyll fluorescence. F<sub>0</sub>, F<sub>m</sub>, Fv/Fm and Y(II) have significantly decreased under high light. The enhancement of the quantum yield of non-regular energy dissipation Y(NO) indicates that both the conversion of photochemical energy and the protective regulating mechanisms are inefficient. The increase of Y(NO) certifies the intensification of dissipation processes which take place mainly in the light harvesting antenna, in phycobilisomes. The high light intensities have led to the photoinhibition of the photosystem (PS) II antenna, affecting the reaction centers because Fv/Fm was decreased. The recovery of PSII photochemical activity was in overall a slow process correlated with light intensity that was used.

**Keywords:** chlorophyll fluorescence; coefficient of photochemical quenching (qP, qL); effective PS II quantum yield ( $Y_{II}$ ); fluorescence yield; minimal fluorescence; maximal fluorescence; maximal PS II quantum yield (Fv/Fm); quantum yield of non-regulated energy dissipation (YNO); recovery.

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#### Introduction

The variability of the excitation spectrum in cyanobacteria is correlated with the adaptation mechanisms of the photosynthetic apparatus with important implications on chlorophyll fluorescence. The photosynthetic apparatus adaptation to various environmental conditions such as light and nutrients (Bryant,1995; Dau, 1994; Demarsac and Houmard,1993; Müller *et al.*,1993; Reuter and Müller,1993) has consequences in changing the phycobilisomes content and the PS I/PS II ratio.

The chlorophyll associated to PS II has a different fluorescence production from PS I (Pfündel, 1998). The chlorophyll amount associated with photosystems is dependent on species, while its variability is induced under stress (Boekema *et al.*, 2001; Falkowski and Raven, 1997; Riethman and Sherman, 1988; Straus, 1994).

The short-time variations of the coupling energy between phycobilisomes and the two photosystems, the so-called transition-states, are associated with the redox state of the electron transport chain. By modulating the energy transport between PS I and PS II (Biggins and Bruce, 1989; Campbell *et al.*, 1998; van Thor *et al.*, 1998; Williams and Allen, 1987) it was defined the chromatic adaptation which is genetically controlled being a phenotypic response to light spectrum. The ratio between phycobilins is changing in relation to the light and its quality (Dubinsky and Stambler, 2009), while the redox state of the photosynthetic components plays an essential role (Campbell and Öquist,1996; Mullineaux and Emlyn-Jones, 2004). This variability of the photosynthetic apparatus is the consequence of an adaptation mechanism, mainly in the peripheral antenna made mostly by phycobilisomes (Bennett and Bogorad, 1973).

The studies accomplished on *Synechococcus* sp. PCC 6803 were focused on the energy transfer from phycobilisomes to photosystems (Krumova *et al.*, 2010; Ma *et al.*, 2008), on the mechanisms for lowering the fluorescence (Bissati *et al.*, 2000), and a protective mechanism of the photosynthetic apparatus in cyanobacteria due to carotenoids was emphasized (Rakhimberdieva *et al.*, 2007; 2007; 2010). In the present study there are presented the results of monitoring the chlorophyll fluorescence in *Synechocystis* sp. PCC 6803 under various light intensities during long periods of time.

#### Material and methods

The cyanobacterium Synechocystis sp. PCC 6803 was grown at room temperature on BG 11 growth medium, under air bubbling and a moderate light intensity of 28  $\mu$ mol m<sup>-2</sup>s<sup>-1</sup> for 9 days. The cells in exponential growth phase were exposed at high light intensities of 2000, 3000 and 4500 µmol m<sup>-2</sup>s<sup>-1</sup>. The light source was an FHI-5000 W fluorescent bulb, while the light intensity was measured with a Quantum Sensor QSPAR Hansatech device. The light-treatment effect was assessed after 15, 30, 45, 60, 75, 90, 105 and 120 minutes. The 120 minutes recovery period since stopping the light treatment was monitored based on fluorescence, which was measured with a Walz Dual-100 fluorometer. The assimilatory pigments (chlorophyll a, carotenoids) were extracted with acetone and they were spectrophotometrically estimated based on specific absorption coefficients (Arnon, 1949; Lichtenthaler and Wellburn, 1983), and their identification was assessed based on the absorption peak with a Jasco V-630 spectrophotometer. The phycobiliproteins were estimated based on Gantt and Lipschultz (Gantt and Lipschulz, 1974). The results are displayed in mg per liter on cell suspension.

#### **Results and discussions**

The *Synechocystis* sp. PCC 6803 culture has reached an optical density  $OD_{680} = 1.567$ . The *in vivo* absorption spectrum has displayed the spectral absorption areas of the photosynthetic apparatus components and the main absorption peaks (fig. 1). Carotenoids (481 nm) and chlorophyll a (441 nm) absorb in the blue spectral area, while phycobilins (628 nm) and chlorophyll a (682 nm) have an absorption peak in the red light.



Fig. 1. *In vivo* absorption spectrum in *Synechocystis* sp. PCC 6803 grown under standard conditions.

The phycobilins are the main pigments of the light-harvesting antenna, while chlorophyll acts in photosystems reaction centers. Through photo-acclimatization the pigments amount is adjusted as an answer to light intensity (Dubinsky and Stambler, 2009). The content of assimilatory pigments which have been identified is displayed in Table 1. The components of the photosynthetic apparatus in cyanobacteria contain chlorophyll a (665 nm) which together with various carotenoids and phycobiliproteins forms the light-harvesting unit. The pigments concentration was measured right before applying the light treatment (control).

**Table 1.** The amount of assimilatory pigments in *Synechocystis* sp. PCC 6803grown under standard conditions (mg/l)

Chlorophyll <u>a</u>	Carotenoids	Phycobiliproteins						
		Phycocyanin	Allophycocyanin	C-phycoerythrin				
6.978	2.052	16.382	4.953	0.335				

According to Rakhimberdieva *et al.* (2004), the major carotenoids in *Synechocystis* are zeaxanthin,  $\beta$ -carotene, echinenone and myxoxatnhophyll which help to energy dissipation through phycobilisomes. The carotenoids play an important role in deactivating the antenna chlorophylls under excess excitation. The protective function of carotenoids is a consequence of their short distance to chlorophylls.



**Fig. 2.** Evolution of the chlorophyll a fluorescence parameters (A) and the quantum yield (B) in *Synechocystis* sp. PCC 6803 under 2000 µmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity.

By exposing the strain *Synechocystis* sp. PCC 6803 to 2000  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensity, the F<sub>0</sub> minimum fluorescence has decreased, excepting the 75 minutes measurement. The decreasing rate at the end of the experiment was 85.82% of the control. The fluorescence main level shows that the Q<sub>A</sub> acceptors are highly oxidized, while the PSII centers are open. The F<sub>m</sub> a maximum fluorescence has decreased together with treatment duration to 56.52% at the experiment end (fig. 2 A). The fluorescence maximum level shows that Q<sub>A</sub> are highly reduced, and the PSII centers are closed. The diminishing of the maximum fluorescence has led to some low values of the variable fluorescence (Fv).

The maximal quantum yield of PSII,  $F_v/F_m$ , has significantly decreased by exposing the cyanobacterial suspensions at 2000 µmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity, generating a rate of final decrease of 51.74% (fig. 2 B). The maximum efficiency shows that light absorbed through PSII is used for reducing Q<sub>A</sub>. The quantum yield of non-regulated energy dissipation Y(NO) has displayed an increase of 152.17%. The effective quantum yield of PSII, Y(II), has proportionally decreased together with light-exposure, showing a decrease rate of 53.44% at the end of the experiment.

The excitation energy regulation induced between the two photosystems is dependent to the moving of phycobilisomes on the thylakoid membrane and which defines the transition state. Two different mechanisms are involved in the state transition induced by light and redox state. The transition state involves the physical movement of phycobilisomes, while the redox transition involves the phycobilisomes movement and the scattering of energy between photosystems (Li *et al.*, 2004).

The photochemical quenching coefficients, qP and qL, have displayed mainly low values when compared to control (fig. 3).



**Fig. 3.** Evolution of the photochemical coefficients (qP, qL) in *Synechocystis* sp. PCC 6803 under 2000 μmol.m<sup>-2</sup>.s<sup>-1</sup> light.

The kinetics of chlorophyll fluorescence induction shows through the analysis of the saturation pulse the evolution of each fluorescence parameter, the duration of oxidation/reduction of the  $Q_A$  primary electron acceptor and also the half time of the fluorescence curve ( $T_{1/2}$ ), which represents the functional size of the antenna. A slow and persistent reduction of the  $Q_A$  primary electron acceptor has occurred, and also a longer period was achieved, certifying a bigger antenna for the control, while in the probe exposed to light the processes were much faster: the maximum fluorescence has decreased, while the  $F_m$  maximum fluorescence was enhanced after illumination (fig. 4).



Fig. 4. The kinetics of chlorophyll fluorescence in *Synechocystis* sp. PCC 6803, control and the variant exposed to 2000 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity, using the saturation pulse method: A – the kinetics of the fluorescence in control sample; B – the induction of fluorescence curve in control; C - the fluorescence kinetics under 2000 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity; D – the fluorescence induction curve at 2000 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity.

The effect of the 3000  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensity on the photochemical activity of PSII is presented in figure 5. The F<sub>0</sub> minimum fluorescence has decreased proportionally to the light exposure period, with 84.21% decrease from the control.

The  $F_m$  maximum fluorescence has significantly decreased with 57.74% at the end of the exposure period. These values have led to the diminishing of the variable fluorescence (Fv) (fig. 5 A). The maximal quantum yield of PSII has progressively decreased during light exposure, with a 57.35%

decrease at the end of the experiment (fig. 5 B). The quantum yield of nonregulated energy dissipation Y(NO) has increased with 146.10%. The effective quantum yield of PSII has decreased proportionally with the exposure period, down to 60.05% at the end of the exposure.



Fig. 5. Evolution of the chlorophyll fluorescence parameters (A) and the quantum yield (B) in *Synechocystis* sp. PCC 6803 under 3000 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity.

The qP and qL photochemical quenching coefficients have displayed variable values under 3000  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light (fig. 6). The oxidation state of the Q<sub>A</sub> primary electron acceptor reflects the difference in the PSII photochemical capacity and also the proportion of the oscillating "openings" during light exposure.



**Fig. 6.** The evolution of the photochemical coefficients (qP, qL) in *Synechocystis* sp. PCC 6803 under 3000  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensity.

The graph of the chlorophyll fluorescence kinetics through the analysis of the saturation pulse shows that a slower and longer reduction of the  $Q_A$  primary electron acceptor has occurred and a longer half time was obtaining, certifying a bigger antenna (fig. 7).



Fig. 7. The kinetics of the chlorophyll fluorescence in *Synechocystis* sp. PCC 6803 under 3000 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity, using the saturation pulse method: A – fluorescence kinetics; B – fluorescence induction curve.

The effect of 4500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensity on the PSII photochemical activity is presented in figure 8. The F<sub>0</sub> minimum fluorescence has gradually decreased during light exposure, with an 81.04% decrease at the end of the test. The F<sub>m</sub> maximum fluorescence has significantly decreased down to 52.54% at the end of the light exposure. The diminishing of the maximum fluorescence has led to the considerably decrease of the variable fluorescence (Fv) (fig. 8 A).

The maximal quantum yield of PSII has proportionally decreased with 49.52% during light exposure (fig. 8 B). The quantum yield of non-regulated energy dissipation Y(NO) has increased with 154.27%. The PSII effective quantum yield (YII) has significantly decreased with 50.05% at the end of light exposure treatment.

The decrease of  $F_0$ ,  $F_m$  fluorescence, and of the effective and maximal quantum yield and also the increase of the quantum yield of the non-regulated energy dissipation Y(NO) certify the inhibition produced by high light on the processes that take place in the photosystems antennas and PSII reaction centers.



Fig. 8. The evolution of the chlorophyll fluorescence parameters (A) and the quantum yield (B) in *Synechocystis* sp. PCC 6803 4500 μmol.m<sup>-2</sup>.s<sup>-1</sup> light.

The photochemical quenching coefficients qP (the fraction of open reaction centers able of pohotochemistry) and qL (oxidized Q<sub>A</sub>), have gain in generally lower values during the 4500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light treatment (fig. 9). qP comes from using the excitation energy inside PSII in order to lead the electron transport from P<sub>680</sub> to Q<sub>A</sub>. "Open" refers to the PSII reaction center where the Q<sub>A</sub> primary acceptor is oxidized and is able of photoreduction, while "closed" denotes the reaction center where Q<sub>A</sub> is reduced and unable to photochemically perform anymore.



Fig. 9. The evolution of the photochemical coefficients in *Synechocystis* sp. PCC 6803 under 4500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light.

The kinetics of chlorophyll fluorescence induction by analyzing the saturation pulse has shown that a slower but longer reduction of the primary electron acceptor  $Q_A$  has occurred, as well as a longer half-time which suggests a bigger antenna (fig. 10).



Fig. 10. The kinetics of the chlorophyll fluorescence in *Synechocystis* sp. PCC 6803 under 4500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensity, using the saturation pulse method: A – fluorescence kinetics; B – fluorescence induction curve.

During the recovery period of the photosynthetic apparatus subsequent to various high light intensities the following results were achieved:  $F_0$  value has remained higher that in control in the samples previously treated with 2000 and 3000 µmol.m<sup>-2</sup>.s<sup>-1</sup> light;  $F_m$  has remained low; the maximal quantum yield has remained low, while the non-regulated energy dissipation has stayed high (fig. 11).



**Fig. 11.** The evolution of the chlorophyll fluorescence parameters in *Synechocystis* sp. PCC 6803 during the recovery period after exposure to various light intensities.

During the recovery period, the photochemical coefficients were significantly reduced (fig. 12). The cells exposed to high light have significantly reduced their photochemical quenching coefficients. The process of recovery of the PSII photochemical functionality was rather slow.



**Fig. 12.** The evolution of the photochemical coefficients in *Synechocystis* sp. PCC 6803 during the recovery period after exposure to various light intensities

The increase of  $F_0$  and decrease of  $F_m$  are due to photoinhibition (van Wijk and Hasselt, 1993). Low  $F_m$  suggests the decrease of the closed  $Q_A$  number, while the interval of the excitation at the marginal receptor is shorter (Laisk *et al.*, 1997). The fluorescence emitting is influenced by the antenna heterogeneity and reduced border heterogeneity (Lazár, 1999). The dissipation of the non-radiative excitation energy in photosystems antenna leads to the  $F_0$  and  $F_m$  decrease and the dissipation of the reaction centers reduce only the  $F_m$  (Gilmore and Yamamoto, 1991).

The PSII photoinhibition is characterized by the quantic efficiency decline, by the decrease in the number of centers involved in the linear transport of the photosynthetic electrons and by increasing the number of centers involved in the excitation energy dissipation (Critchley and Russell, 1994). Vass *et al.* (1992) have observed that and increased  $F_0$  may be the consequence of the double reduction and of the putative protonation of  $Q_A$ , or is it is due to the separation of the PSII reaction centers (Srivastava *et al.*, 1997).

The decrease of Fv/Fm due to the increase of  $F_0$  or decrease of  $F_m$  is a conventional measure of photoinhibition as well as an indicator of the PSII photochemical efficiency reduction because the Fv/Fm ratio is strongly connected to the quantum yield of the photosynthetic  $O_2$  evolution or  $CO_2$ assimilation. A fast increasing of  $F_0$  may be induced by inactivating of a part of the PSII reaction centers (Harel *et al.*, 2004). The PSII photoinactivation (photoinhibition), the recovery reaction and the interactive photoprotective mechanisms are fundamental components of metabolism, physiology and ecophysiology of oxygenic photoautotrophic organisms (Campbell and Tyystjärvi, 2012).

#### Conclusions

*Synechocystis* sp. strain PCC 6803 has displayed an increase and a composition of the assimilatory pigments specific for its species.

Under high light intensities, the chlorophyll fluorescence parameters have significantly decreased in the illuminated samples ( $F_0$ ,  $F_m$ , Fv/Fm and YII). The low values certify the use of absorbed energy mainly as chemical energy through photochemical charges separation in the RC II reaction centers. The high value of the quantum yield of the non-regulated energy dissipation shows that both the photochemical energy conversion and the protective regulator mechanisms are inefficient. The increase of the quantum yield of non-regulated energy dissipation certifies the intensity of the dissipation processes that occur mainly in the light harvesting antenna, in the phycobilisomes.

The high light intensities that were used in this study have led to photoinhibition of the PSII antenna, affecting the reaction centers because the maximal quantum yield has decreased. The efficiency of the energetic mechanisms in the antenna and reaction centers is observed with accuracy through the kinetics of chlorophyll fluorescence induction by analyzing the saturation pulse.

The recovery of the PSII photochemical activity was, in general, a slow process, in relation with the light intensity that the cell suspensions were exposed to.

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# THE STUDY OF PHOTOSYSTEM II (PS II) ACTIVITY BASED ON CHLOROPHYLL FLUORESCENCE IN APHANIZOMENON ELENKINII AICB 709, UNDER LIGHT STRESS CONDITIONS

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SUMMARY. High light-induced changes of chlorophyll fluorescence during 2 hours are analyzed in cyanobacteria Aphanizomenon elenkinii AICB 709. The light of 800 µmol.m<sup>-2</sup>.s<sup>-1</sup> produced the increasing of F0 and Fm in the first part of light exposure. Fv and the maximal and effective quantum yields decreased after 30 min. of light exposure. The opening state and the ratio of the reaction centers decreased. The light of 1500 µmol.m<sup>-2</sup>.s<sup>-1</sup> induced the rising of F0, Fm and Fv and the decreasing of maximal and effective quantum yields, relative to control. The decreasing of photochemical coefficients means that the fraction of the opened reaction centers and their ratio also were significantly diminished. The light of 2100 umol.m<sup>-2</sup>.s<sup>-1</sup> determined the increasing of F0 in the first 15 min, and the decreasing of Fm, Fv, quantum yields and photochemical coefficients. The quantum yield of non-regulated energy dissipation Y(NO) enhanced in every light intensity that was used. Changes in fluorescence chlorophyll are caused by structural changes in the antenna and the PS II reaction centers due to photoinhibition produced by high light. The fast recovery of the photosynthetic parameters was obtained in the samples exposed to high light intensity.

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**Keywords:** carotenoid, chlorophyll <u>a</u>, coefficient of photochemical quenching, effective quantum yield, minimal and maximal fluorescence, maximal quantum yield, phycobilisomes, quantum yield of non-regulated energy dissipation, recovery, saturation pulse method.

# Introduction

The large density of the photons flux causes damages in the photosynthetic apparatus due to the excessive absorbed energy that can not be used in photochemistry. To avoid the photo-oxidative damages, the photosynthetic microorganisms developed regulating mechanisms for the efficient conversion of the excitation energy into heat (Jahns and Miehe, 1996).

The inactivation process is named photoinhibition. The inactivation level of PS II was quantified through the determination of slow relaxing component (qI) of the fluorescence non-photochemical quenching (qN) (Krause and Weis, 1991; Quick and Stitt, 1989) and through the measuring of the changes in Fv/Fm.

Photoinhibition involves at least two inactivation stages. The first stage is considered reversible and is produced in approximately one hour without any damages of PS II (Leitsch *et al.*, 1994). The second stage is supposed to be associated with damages of the PS II reaction center. It is likely that this stage is oxygen dependent and it is reversible through the replacement of D1 protein (Aro *et al.*, 1993; Leitsch *et al.*, 1994).

Photoinhibition is a process common to plants and cyanobacteria (Krupa *et al.*, 1990; Lönneborg *et al.*, 1988; Samuelsson *et al.*, 1985; 1987). The photoinhibition installs when the captured energy exceeds the dissipated energy as a result of the rising in light intensity or as a result of moderate light when an excessive harvesting antenna is present, or if the low temperature limits the dissipation of the energy (Huner *et al.*, 1993). 36

The most sensitive spot to photoinhibition is the reaction center of PS II through which the key reactions become inactive in the first place, *in vivo*. *In vitro*, photoinhibition occurs after the primary charge separation if the electron donation to the oxidized reaction center of PS II is blocked, or instead through over-reducing of the primary acceptor  $Q_A$ , on the electron accepting side of PS II (De Las Rivas *et al., 1992*). Both types of inhibition are the result of the degradation of D1 protein (Shipton and Barber, 1991; Vass *et al., 1992*). On the whole, the photosystem II photoinhibition is characterized by the declining in the quantum efficiency, number of reaction centers that took part to the linear photosynthetic electron transfer and the increasing in the number of centers which are involved in the dissipation of the light intensity is excessive and may cause damages at the level of PS II reaction centers (Casper-Lindley and Björkman, 1998; Long*et al., 1994*).

Changes in light intensity or in temperature induce an imbalance between the energy of the light absorbed through photochemistry and the energy used in metabolism. This type of energetic imbalance is pointed out by the shift of excitation pressure of PS II photosystem, which reflects the reductive relative state of the photosystem (Huner *et al.*, 1998).

The light harvesting regulation is necessary to balance the absorption and the usage of the light energy and to minimize the photo-oxidative damaging potential (Müller *et al.*, 2001). The great variety of cyanobacterial photosynthetic apparatus is the consequence of an adaptation mechanism located in the peripheral antenna formed by phycobilisomes (Bennett and Bogorad, 1973).

The results of the present study refer to the photosystem PS II activity pointed out by the chlorophyll fluorescence measurements in the cyanobacteria *Aphanizomenon elenkinii* AICB 709, exposed to different light intensities for prolonged periods of time.

### Materials and methods

The cyanobacteria *Aphanizomenon elenkinii* AICB 709 (Nostocales) was cultivated at room temperature on BG 11 medium, using a medium light intensity of 26  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup>, in air-lift conditions. The growth period was set to 8 days. The cells found in exponential growth rate were subjected to high light intensities of 800, 1500 and 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup>, for a two hour period. The light source was an incandescent lamp FHI-5000 W and the light intensity was measured with Quantum Sensor QSPAR Hansatech.

The effect of light exposure was measured during 15, 30, 45, 60, 75, 90, 105 and 120 min. During the recovery period, 120 min. from ceasing the light stress, specific fluorescence measurements were taken. The photosystem II activity was evaluated by chlorophyll fluorescence parameters of light exposed probes, using a Waltz Dual-100 fluorometer.

The assimilatory pigments (chlorophyll <u>a</u> and carotenoids) were extracted in acetone and spectrophotometrically estimated based on specific absorption coefficients (Arnon, 1949; Lichtenthaler and Wellburn, 1983), thus their identification was done based on the maximal absorption peak using a Jasco V-630 spectrophotometer. Phycobiliproteins were estimated after Gantt and Lipschultz (1974). The results were expressed in mg/l cellular suspension.

#### **Results and discussions**

The cell suspension of *Aphanizomenon elenkinii* AICB 709 showed on optical density  $OD_{680}$ =1.080. The *in vivo* absorption spectra emphasized the blue spectra where the carotenoids (483 nm) and chlorophyll a (438 nm) absorbed energy and the red spectra where the phycobilins (638 nm) and chlorophyll a (680 nm) absorbed energy (fig. 1). The pigment composition is listed in Table 1.



Fig. 1. *In vivo* absorption spectra of *Aphanizomenon elenkinii* AICB 709 cultures, maintained in standard growth conditions.

**Table 1.** The quantities of assimilatory pigments inAphanizomenon lenkinii AICB 709 cultures, maintainedin standard growth conditions (mg/l).

Chlorophyll	Carotenoids	a/c	Phycobiliproteins				
<u>a</u>			Phycocyanin	Allophycocyanin	C-phycoerythrin		
5.592	1.357	4.12	4.755	6.086	4.394		

In cyanobacteria, all chlorophylls are located in the antenna nucleus of PS I and PS II: the chlorophyll/ $P_{700}$  ratio is 100, and that of chlorophyll/ $P_{680}$  is 40 (Karapetyan, 2007).

Due to the exposure of *A. elenkinii* AICB 709 cells to different light intensities, the photochemical activity of the photosynthetic apparatus suffered radical changes. The effect of 800  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> resulted in a significant rising of the minimal fluorescence F<sub>0</sub> between 45 and 90 min. reaching a maximal increase of 178.65% at 75 min. The maximal fluorescence F<sub>m</sub> maintained above control values for 60 min. and then decreased to 40.8% in the end. F<sub>v</sub> decreased significantly after the first 30 min of light exposure (fig. 2 A). The maximal quantum yield Fv/Fm of PSII photosystem (photochemical efficiency) decreased to 0.35% after 30 min. in the end of the exposure, relative to control (fig. 2 B). The quantum yield of the electron transfer to

PSII (photochemistry) in the presence of light is the measure of the whole efficiency of the reaction centers and asserts the ratio of the absorbed energy by the antenna which is then used in photochemistry. The effective quantum yield  $Y_{(II)}$  begun to decrease slowly after 30 min. reaching 2.6% at the end. The quantum yield of non-regulated energy dissipation Y(NO) significantly increased after 45 min. reaching a finale increase of 196.6 % to the end of the exposure. Based on the recorded values of quantum vield, after 30 min. of light exposure, some structural changes occurred in the antenna, which describe a lack of energy in the thylakoid membrane. The ratio of the opened reaction centers of PSII was high (q<sub>P</sub>) and the non-photochemical excitation energy dissipation was insignificant. All the fluorescence parameters values emphasized the inactive state of the photosystem PSII antenna and the lowering of the photosynthetic performance. Non-radiative dissipation of the excitation energy in the photosystems antenna produced the diminishing of F<sub>0</sub> and F<sub>m</sub> and the dissipation from the reaction centers reduces only Fm (Gilmore and Yamamoto, 1991). The fluorescence diminishing associated with the photoinhibition is not directly linked with the degradation of the PSII reaction centers (Briantais et al., 1988).



**Fig. 2.** The effect of 800 μmol.m<sup>-2</sup>.s<sup>-1</sup> light exposure on the evolution of chlorophyll fluorescence parameters (A) and the quantum yields (B) in *Aphanizomenon elenkinii* AICB 709.

On the whole, the photochemical coefficients qP and qL reached values below 1 during the entire period of high light exposure. The significant decreasing was obtained at the end of the exposure, being 87.4% relative to control. The opening level of the reaction centers and their ratio decreased relative to control due to the exposure at 800µmol.m<sup>-2</sup>.s<sup>-1</sup> (fig. 3). After 90 min. of light exposure, maximal values for non-photochemical energy dissipation coefficient qN was recorded. The persistence of the minimal excitation pressure (1-qP) asserts the existence of a dimensioned antenna in the context of the reduction of photosynthetic efficiency.



**Fig. 3.** The effect of 800 μmol.m<sup>-2</sup>.s<sup>-1</sup> light exposure on the evolution of photochemical coefficients in *Aphanizomenon elenkinii* AICB 709.

Based on the chlorophyll fluorescence induction kinetics throughout the saturation pulse the intensity of each parameter may be estimated: the duration of oxidation/reduction of the primary electron acceptor  $Q_A$  and the half time of the fluorescence curve ( $T_{1/2}$ ) which expresses the functional size of the antenna: a rapid reduction of  $Q_A$  decreases the doubling time; a slow reduction of  $Q_A$  increases the doubling time (fig. 4).



Fig. 4. The effect of high light exposure on the kinetics of chlorophyll fluorescence based on the saturation pulse method in *Aphanizomenon elenkinii* AICB 709:

- A- the chlorophyll fluorescence kinetics in control;
- B the fluorescence induction curve in control;
- C- the fluorescence kinetics at 800 µmol.m<sup>-2</sup>.s<sup>-1</sup>;
- D- the fluorescence induction curve at 800  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup>.

The light effect at 1500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> over *A. elenkinii* AICB 709 cell suspension is showed in Figure 5. F<sub>0</sub> increased in the first 75 min., than it decreased to 85%, relative to control. The maximal fluorescence F<sub>m</sub> lowered proportional with the exposure time reaching 41,9% at the end, thus the closure of the reaction center is much diminished. Fv decreased significantly as a result of light exposure (fig. 5 A).

The PSII maximal quantum yield  $F_v/F_m$  decreased significantly to 0.52% at the end of the exposure and the effective quantum yield Y(II) decreased to 3% (fig. 5 B). The quantum yield of non-regulated energy dissipation, Y(NO), reached 196% at the end of the exposure.



**Fig. 5.** The effect of 1500 μmol.m<sup>-2</sup>.s<sup>-1</sup> light exposure on the evolution of chlorophyll fluorescence parameters (A) and on the quantum yields (B) in *Aphanizomenon elenkinii* AICB 709.

The photochemical coefficients qP and qL significantly lowered after 45 min of exposure and they reached maximal values towards the end of the exposure when the coefficient of non-photochemical energy dissipation qN was activated. The diminishing of photochemical coefficients indicated that the ratio of the opened reaction centers and their ratio also lowered significantly (fig. 6).



**Fig. 6.** The effect of 1500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light exposure on the evolution of photochemical coefficients (qP, qL) in *Aphanizomenon elenkinii* AICB 709.

The fluorescence curve, the oxidation/reduction time of the primary electron acceptor  $Q_A$  and the half time of the fluorescence curve  $(T_{1/2})$  which expresses the functional size of the antenna can be analyzed based of the chlorophyll fluorescence induction kinetics through the analysis of the saturation pulse. A quick transition of the primary electron acceptor  $Q_A$  which defines a small size antenna has occurred (fig. 7).



**Fig. 7.** The chlorophyll fluorescence kinetics using the saturation pulse method in *Aphanizomenon elenkinii* AICB 709 exposed to 1500 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity. A- the chlorophyll fluorescence kinetics; B- the fluorescence induction curve.

The effect of 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensity on PSII photochemistry is shown in Figure 8. The minimal fluorescence F<sub>0</sub> increased after 15 min of light exposure, reaching a maximal value of 305.9% at 45 min, thus the primary electron acceptor Q<sub>A</sub> becomes oxidized and the reaction centers are opened, capable of photochemical performance in reducing Q<sub>A</sub>. The maximal fluorescence F<sub>m</sub> decreased after 60 min of light exposure leading to a finale decrease of 48.8%, relative to control. Fv decreased significantly due to light exposure (fig. 8, A).

The maximal quantum yield  $F_v/F_m$  of PS II decreased significantly to 0.29% at the end of the exposure, relative to control. The effective quantum yield  $Y_{(II)}$  of PSII, decreased significantly reaching 2.2% toward the end of the exposure. The quantum yield of non-regulated energy dissipation Y(NO) increased 198.2% at the end of light exposure (fig. 8, B).



**Fig. 8.** The effect of 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light exposure on the evolution of chlorophyll fluorescence parameters (A) and on the quantum yields (B) in *Aphanizomenon elenkinii* AICB 709.

The diminished values of photochemical efficiency  $F_v/F_m$  and the reduction of the quantum yields  $Y_{(II)}$  emphasize the initiation of photosynthesis down regulation process. The mechanism of down regulation creates energy dissipation centers leading to a homogeneous down regulation through the

increasing of the quantum yield of non-regulated energy dissipation Y(NO). The mechanism of photosystems antenna is involved in dissipating the down regulation centers and the D1 turnover is important in the initiation and function of down regulation (Critchley and Russell, 1994).

The coefficients of photochemical quenching qP and qL decreased significantly at 45 and 90 min. of light exposure (fig. 9). After 45 min. of light exposure, the coefficient of non-photochemical energy dissipation qN was activated.



**Fig. 9.** The effect of 2100 μmol.m<sup>-2</sup>.s<sup>-1</sup> light exposure on the evolution of photochemical coefficients in *Aphanizomenon elenkinii* AICB 709.

Through the analysis of the saturation pulse, some changes in the fluorescence curve are visible, on chlorophyll fluorescence kinetics chart in the means of the oxidation/reduction period of the primary electron acceptor  $Q_A$  and in the half time of the fluorescence curve  $(T_{1/2})$ . A prolonged reduction of the primary electron acceptor  $Q_A$  has produced in a very short time, which asserts the existence of a diminished antenna (fig. 10).



**Fig. 10**. The chlorophyll fluorescence kinetics using the saturation pulse method in *Aphanizomenon elenkinii* AICB 709 exposed to 2100 μmol.m<sup>-2</sup>.s<sup>-1</sup> light intensity. A- the chlorophyll fluorescence kinetics; B- the fluorescence induction curve.

During the recovery of the photosynthetic apparatus, the minimal fluorescence remained high relative to control (fig. 11). The maximal fluorescence increased to 67% in the case of 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light exposing. The PSII maximal quantum yield and the effective quantum yield Y<sub>(II)</sub> remained diminished, relative to control. The quantum yield of non-regulated energy dissipation Y(NO) remained high during the recovery period, quick recoveries being recorded at 1500 and 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light intensities. The photochemical coefficients reached slightly diminished values relative to control. On the whole, a more rapid recovery was observed in the cyanobacterial cell suspensions exposed to 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> and 1500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> respectively.



**Fig. 11.** The effect of various light exposures on the evolution of chlorophyll fluorescence parameters (A) and on the quantum yields (B) in *Aphanizomenon elenkinii* AICB 709, in the recovery period.

## Conclusions

The *Aphanizomenon elenkinii* AICB 709 strain was cultivated at room temperature on BG 11 medium, using a medium light intensity of 26  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup>, in air-lift conditions. The growth curve and the composition in assimilatory pigments were species-specific. The components of the photosynthetic apparatus comprise chlorophyll <u>a</u> which together with various types of carotenoids and phycobiliproteins forms the light harvesting antenna.

The  $F_0$  increased between 45 and 90 min. in conditions of 800  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light exposure.  $F_m$  increased in the first 60 min. and than decreased to 40.8% at the end. Fv significantly decreased after the first 30 min. of light exposure. The maximal quantum yield Fv/Fm decreased in the first 30 min, reaching 0.35% at the end of the exposure. The effective quantum yield Y<sub>(II)</sub> gradually decreased after 30 min. reaching 2.6% at the end of light exposure. The quantum yield of non-regulated energy dissipation, Y(NO), increased significantly after 45 min. The ratio of the opened reaction centers of PS II was high and the non-photochemical dissipation of the excitation energy was negligible. The fluorescence parameters values emphasize the inactivation state of the PSII photosystem antenna and the decreasing of 48

the photosynthetic performances. The photochemical coefficients qP and qL decreased, thus asserting that the opening state of the reaction center and their ratio diminished also.

The  $F_0$  increased in the first 75 min. than decreased to 83 % in conditions of 1500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> light exposure.  $F_m$  decreased to 41.9% in the end, which certifies that the closure of the reaction centers is much diminished.

The maximal quantum yield decreased to 0.52 % at the end of the exposure and the effective quantum yield decreased to 3%. The quantum yield of non-regulated energy dissipation recorded on increasing of 196 % at the end of light exposure. The photochemical coefficients decreased, and toward the end they reached maximal values.

 $F_0$  increased after 15 min. in conditions of 2100 µmol.m<sup>-2</sup>.s<sup>-1</sup> light exposure, thus the primary acceptor Q<sub>A</sub> become oxidized and the reaction centers are opened, capable of photochemical performance.  $F_m$  decreased after 60 min of light exposure. The maximal quantum yield  $F_v/F_m$  of PSII significantly decreased to 0.29% relative to control at the end of light exposure. The effective quantum yield Y<sub>(II)</sub> of PSII decreased significantly reaching 2.2 % at the end of light exposure. The quantum yield on nonregulated energy dissipation Y(NO) recorded an increasing of 198.2 % at the end of the experiment. The photochemical coefficients significantly decreased at 45 and 90 min. of light exposure.

Through the analysis of the saturation pulse, some changes in the fluorescence curve are visible on chlorophyll fluorescence kinetics chart in the means of the oxidation/reduction period of the primary electron acceptor  $Q_A$  and in the half time of the fluorescence curve ( $T_{1/2}$ ). A prolonged reduction of the primary electron acceptor  $Q_A$  has produced in a very short time, which asserts the existence of a diminished antenna.

A faster recovery of the fluorescence parameters was observed in the cyanobacteria cell suspensions exposed to 2100  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> and 1500  $\mu$ mol.m<sup>-2</sup>.s<sup>-1</sup> respectively. This cyanobacteria strain proved to be very sensitive to the light intensity.

The low values of the photochemical efficiency and that of the quantum yield and the dissipation increments emphasized the initiation of the photosynthesis photoinhibition process, leading to changes in the antenna and the reaction centers, influencing the photochemistry and the fluorescence emission.

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# AN ESTIMATION FOR LOWER BOUND OF p53 IN DNA HEALING PROCESS; A MATHEMATICAL APPROACH

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**SUMMARY:** The main goal here is to estimate a lower bound for the level of the p53 protein which is necessary for the DNA damage is repaired by a coeffective work of activated p53 and ATM-p. We consider a system of ODEs as the mathematical model of interactions between the proteins p53, Mdm2, and ATM-p. The model is biologically motivated and can simulate effects of proteins on the DNA damage suppression. It is shown that positive amounts of ATM-p may force the system to pass through a saddle node bifurcation which in turn indicates the lower bound for p53. We see that if the amount of p53 is less that the lower bound then the DNA damage is not suppressed. Finally we use the bifurcation equation and show that how the Hill coefficient of p53 must be controlled in order to optimize the healing process.

**Keywords:** p53-Mdm2 oscillatory model, ATM-p protein, Saddle node bifurcation, Center Manifold, Hill constant.

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## Introduction

In the last decades, biologists have been attempting to treatment the cancer and bind developing tumors. Nowadays, we know that a protein which is called p53 has an effective role in tumor suppressing such that, its disruption is associated with approximately 50 to 55 percents of human cancers (Almong and Rotter, 1997; Bar-Or *et al.*, 1999).

By lab observations, biologists consider three major functions for p53: as it arrests the cell cycle and gets enough time to correct the DNA damage, has an indirect role in transcription of DNA corrective proteins and finally if the above methods are not effective, performs apoptosis (programmed cell death). Because some of the cellular effects of activated p53 can be irreversible, keeping p53 function under tight control in normal cells is critical. A key player in the regulation of p53 is the Mdm2 protein. Also p53 is a transcription factor of proteins which regulates Mdm2 level (Almong et al., 2000; Sigal and Rotter, 2000). This interaction imposes an intercellular negative feedback loop, which probably serves to keep p53 in tight check and to rapidly terminate the p53 response once a p53-activating stress signal has been effectively deal with (Bar-Or et al., 2000). In fact, once a DNA damage occurs, an intermediate protein which is called ATM is activated, which in turn activates p53 and Mdm2 by phosphorylating this proteins (Kohn and Pommier 2005; Banin et al., 1998). As there is a direct correspondence between the works of p53 and its level, so it is important to realize the dynamics due to the co-effects of proteins involved in this intercellular process.

In the last decade, mathematical models with capability to predict behaviors and simulate dependence of dynamics on parameters and variables provide a significant improvement in the dynamics analysis.

There are many mathematical models simulating the reaction between interfered proteins in cell cycle and predict the effect of parameters on final behavior of these proteins (for example see McCoy *et al.*, 2003; Hinow *et al.*, 2006; Qi *et al.*, 2007; Wagner *et al.*, 2005; Ciliberto *et al.*, 2005; Yan and Zhuo, 2006 and references within).

A recent elementary mathematical model which is motivated biologically, simulates this interaction as below (Chickarmane *et al.*, 2007).

$$\begin{cases} x' = \alpha_0 + \alpha_1 \frac{x^n}{k_1 + x^n} - \gamma_1 x y - \gamma_2 x \\ y' = \alpha_2 + \alpha_3 \frac{x^4}{k_2 + x^4} - \gamma_3 y. \end{cases}$$
(1)

Where x(t) and y(t) are the level of p53 and Mdm2, respectively. In the first equation  $\alpha_0$  represents the production rate of p53 and Mdm2, in fact it is the phosphorylation rate of p53 by ATM-p, second term represents an autocatalytic process which we assume exists due to positive feedback of p53 on itself and is described with a Hill coefficient n, which is a chemical index depends on protein molecular structure, third term describes decreasing the level of p53 by Mdm2 and the fourth term represents the degradation of p53 independently of Mdm2. Likewise, in the second equation,  $\alpha_2$  describes a basal production level of Mdm2, and the second term represents the activation of Mdm2 by p53, described by a Hill equation with exponent 4, the last term represents the degradation of active Mdm2.

Experimental observations show that the DNA damage activates phosphorylation process of ATM which in turn switches on the p53-Mdm2 oscillatory behavior. When the damage is repaired this oscillation is switched off by ATM-p. This is in spirit of Ciliberto, Novak and Tyson (Ciliberto *et al.*, 2005) and can be found through numerical plots of (1). This is the significance of the model (Chickarmane *et al.*, 2007) which makes it biologically motivated. As ATM-p has an important role in activating of p53, ATM-p has a direct role in  $\alpha_0$ . Thus, in the simplest form, by assuming that this effect works linearly, the first equation in (1) must be replaced by

$$x' = z + \alpha_1 \frac{x^n}{k + x^n} - \gamma_1 x y - \gamma_2 x.$$

Here z(t) stands for the amounts of the ATM-p protein at time t, which holds in the following equation

$$z' = \alpha_{1s} \frac{z(\mu - z)}{k_{1s} + \mu - z} + \alpha_{2s} \frac{z}{(k_{0d} + w)(k_{2s} + z)},$$

where w simulates the DNA damage signal. The first term of the above equation, with coefficient  $\alpha_{1s}$  and the second term, with coefficient  $\alpha_{2s}$ show phosphorylation and dephosphorylation rates of the ATM protein, respectively. The parameter  $k_{1s}$  controls dependence of phosphorylation rate of the ATM protein to the level of ATM. Similarly, the parameters  $k_{2s}$  and  $k_{0d}$  control dependence of dephosphorylation rate of the ATM protein to the level of ATM-p and damage signal, respectively.

Finally, the DNA is repaired by the combined actions of ATM-P and p53, which is simulated as the following equation

$$w' = -\alpha_d w x z$$

where  $\alpha_d$  is the rate of repair in dimensionless unites. Therefore the simultaneous model of reaction between levels of these proteins gathers in the following system of differential equations

$$x' = z + \alpha_1 \frac{x^n}{k_1 + x^n} - \gamma_1 xy - \gamma_2 x$$
  

$$y' = \alpha_2 + \alpha_3 \frac{x^4}{k_2 + x^4} - \gamma_3 y$$
  

$$w' = -\alpha_d wxz$$
  

$$z' = \alpha_{1s} \frac{z(\mu - z)}{k_{1s} + \mu - z} + \alpha_{2s} \frac{z}{(k_{0d} + w)(k_{2s} + z)}.$$
(2)

56

As soon as damage occurs, ATM phosphorylates itself which activates the repairing process and damps the damage signal in conditions. In fact, RabieiMotlagh and Afsharnezhad specify an area in the parameter space in which ATM-p suppresses DNA damage by a co-effective work simulated in two last equations of the model (2) (Rabieimotlagh and Afsharnezhad, 2010). They suppose that

$$\Lambda = \mu(\mu + k_{1s}) - k_{1s}k_{2s}, \ U = k_{1s}(\mu + k_{1s} + k_{2s}) \text{ and define}$$
  
$$\Delta = (\mu + k_{1s})\alpha_{2s} - \mu k_{0d}k_{2s}\alpha_{1s}, \ \Omega = \alpha_{2s}\sqrt{U} / \left(\alpha_{1s}(U - \sqrt{U})(\sqrt{U} - k_{1s})\right) - k_{0d}.$$

They proved that change in the sign of  $\Delta$  and  $\Omega$  may cause a bifurcation in the behaviour of the damage signal, (see theorem 2, Rabieimotlagh and Afsharnezhad, 2010) which can biologically interpret the DNA healing process and permanent damage. Furthermore, if the sufficient conditions for the DNA healing process satisfy, except in a null set of this area, always level of ATM-p should be eventually nonzero (see cases (c),(d) and (f) of table A of Rabieimotlagh and Afsharnezhad, 2010). This means that, in order to have the cancer disease suppressed, any control on the co-effective work of ATM-p and p53 must guarantees that the amount of the ATM-p protein remains eventually nonzero.

In what follows, we consider the two first equations of the model (2) and investigate sufficient conditions that the DNA healing process is successfully accomplished. Indeed from the first equation of (2) we know that if the amount of ATM-p is eventually nonzero, then the amount of activated p53 should be also eventually nonzero. Thus we must indicate the lower bound for the amount of activated p53 for which the DNA damage is repaired eventually by co-effective work of ATM-p and p53. We see that the non zero amount of ATM-p causes a saddle-node bifurcation and the stable manifold of the saddle point divides the phase space into two distinct regions. The right hand side region provides the sufficient conditions (the lower bound for amount of p53) for the DNA healing process. Furthermore,

we obtain the bifurcation equation with respect to the amount of ATM-p, the basal production level of Mdm2 and the Hill coefficient  $k_1$ . This helps us to understand the co-effect of the ATM-p and p53 on the healing process; indeed, as soon as the amount of ATM-p remains nonzero, the bifurcation occurs. This, in turn, controls the amount of p53 such that the damage signal is eventually damped. In the last part of the paper we put a control on the amounts of ATM-p in order to optimize the damping process.

#### p53-Mdm2 Oscillatory Model

Let consider the oscillatory model of p53 and Mdm2 as the following system  $% \left( {{{\rm{S}}_{\rm{s}}}} \right)$ 

$$\begin{cases} x' = z + x(\alpha_1 \frac{x^{n-1}}{k + \beta + x^n} - \gamma_1 y - \gamma_2) = 0\\ y' = \theta + \alpha_3 \frac{x^4}{k + x^4} - \gamma_3 y = 0. \end{cases}$$
(3)

Here z,  $\beta$  and  $\theta$  are parameters of system. Therefore we can consider this system as a perturbation with respect to the z,  $\beta$  and  $\theta$  of the system (1). The first step is to prove the following theorem.

**Theorem 1.** For  $\theta = z = \beta = 0$ , the system (3) has a nonhyperbolic critical point, say  $(x_1, y_1)$ , which is a bifurcation point. In fact, in the  $\beta = 0$  plane the system has a saddle node bifurcation and in z = 0 plane it has a transecritical bifurcation

For proving theorem 1, let define unperturbed maps

$$f(x) = \frac{1}{\gamma_1} (\alpha_1 \frac{x^{n-1}}{k+x^n}) - \gamma_2 \quad , \quad g(x) = \frac{\alpha_3}{\gamma_3} (\frac{x^4}{k+x^4}).$$

Notice that the unperturbed system (3) (i.e. when  $\theta = z = \beta = 0$ ) has the origin as a critical point with the linear part  $\begin{pmatrix} -\gamma_2 & 0 \\ 0 & -\gamma_3 \end{pmatrix}$  thus for z,  $\beta$ and  $\theta$  sufficiently small, the origin is a hyperbolic sink. Other steady states are obtained by x' = 0, y' = 0 which imply respectively

$$y = f(x) = \frac{1}{\gamma_1} (\alpha_1 \frac{x^{n-1}}{k + x^n}) - \gamma_2$$
,  $y = g(x) = \frac{\alpha_3}{\gamma_3} (\frac{x^4}{k + x^4})$ 

The points, at which the graphs of f and g meet each other, are the singularities of the unperturbed system. For study of possibility of existence of this points notice that f has a maximum in  $x_0 = \sqrt[q]{k(n-1)} > 0$ . As  $f(0) = -\gamma_2 / \gamma_1$  and g(0) = 0 so if  $f(x_0) < g(x_0)$  then the graph of f dose not crosses the graph of g for x > 0. Also if  $f(x_0) > g(x_0)$  then these two graphs have two cross points for x > 0. Thus by considering parameter  $v = f(x_0) - g(x_0)$  we have proved the following lemma.

**Lemma 3.** In increasing the value of v from negative values to positive ones, there is a unique value  $v_0 \ge 0$  such that:

1) For  $v < v_0$  system (3) has no critical point for x > 0 and for  $v = v_0$ the system (3) has one critical point  $(x_1, y_1)$  with  $0 < x_1 < x_0$ .

2) For  $v > v_0$  the system (3) has exactly two critical points in the x > 0 area. Thus for  $v = v_0$  a saddle node bifurcation occurs at  $(x_1, y_1)$ ; (see Fig. 1).

As the saddle node bifurcation may permanent with respect to small perturbations (Guckenheimer and Holmes, 1993), we show that for  $|\theta|, |\beta|, |z| > 0$  sufficiently small, system (3) has this type of bifurcation; however it is not necessarily the only bifurcation which occurs. We will see latter that the system has also a trasecritical bifurcation. In the next section we apply the center manifold method and compute the bifurcation equation.



**Figure 1.** A saddle node bifurcation occurs for  $v = v_0$  at  $(x_1, y_1)$ .

## Center manifold method

Let  $(x_1, y_1) = (a, f(a))$ , with  $0 < x_1 \le x_0$ , be the tangent point of the graphs (see case  $v = v_0$  in (Fig. 1)). The linear part of the system at  $(x_1, y_1)$  is given by

$$\begin{pmatrix} F_{x}(x_{1}, y_{1}) & F_{y}(x_{1}, y_{1}) \\ G_{x}(x_{1}, y_{1}) & G_{y}(x_{1}, y_{1}) \end{pmatrix}$$

where  $F(x, y) = \gamma_1(xf(x) - xy), G(x, y) = \gamma_3(g(x) - y).$ 

Since f'(a) - g'(a) = 0 so two eigenvalues of the system are given by  $\lambda_1 = 0$  and  $\lambda_2 = a\gamma_1 f'(a) - \gamma_3$ . Since  $f'(a) = g'(a) = (\frac{\alpha_3}{\gamma_3}) 4a^3 k / (k + a^4)^2$  so  $\lambda_2 \ge 0$  if and only if  $Aa^8 + (A-2)2k a^4 + Ak^2 \le 0$ , where  $A = \gamma_3^2 / \alpha_3 \gamma_1$ .

But  $Aa^8 + (A-2)2k a^4 + Ak^2 = 0$  if and only if  $a^4 = (k (2-A) \pm 2k \sqrt{1-A})/A$ . This implies that  $A \le 1$ , but in this case  $A-2 \le 0$  and so  $a^4 < 0$  which is impossible. This guarantees that  $\lambda_2 = a\gamma_1 f'(a) - \gamma_3 < 0$ . Thus, (a, f(a)) has a one dimensional center manifold and a one dimensional stable manifold. This means that the saddle node bifurcation causes two hyperbolic steady states of saddle type and sink type respectively. Furthermore the bifurcation equation is a one dimensional equation. For finding this

equation, we change the system to its standard form and calculate the associated one dimensional center manifold. Whoever, we omit the long calculations and give the results. At first, we transfer  $(x_1, y_1) = (a, f(a))$  to the origin and obtain

$$\begin{cases} x' = F(x, y, z) \\ y' = G(x, y, \theta) \\ z' = 0 \\ \theta' = 0 \\ \theta' = 0 \end{cases}$$
(4)

where

$$F(x, y, z) = z + \gamma_1(x+a)(f(x+a, \beta) - f(a, 0) - y),$$
  

$$G(x, y, \theta) = \theta + \gamma_3(g(x+a) - g(a) - y),$$

with

$$f(x,\beta) = (\alpha_1 x^{n-1} / (k+\beta+x^n) - \gamma_2) / \gamma_1,$$
  
$$g(x) = \alpha_3 x^4 / \gamma_3 (k+x^4).$$

Notice that the Jordan form of the system (4) at the origin is of the form

	$(\lambda_2)$	0	0	0	0)
	0	0	1	0	0
J =	0	0	0	0	0
	0	0	0	0	0
	0	0	0	0	0)

Thus, the standard form of system is as the following

$$\begin{aligned} x' &= \lambda_2 x + H_1(x, y, z, \beta, \alpha_2) \\ y' &= z + H_2(x, y, z, \beta, \alpha_2) \\ z' &= 0 \\ \beta' &= 0 \\ \theta' &= 0, \end{aligned} \tag{5}$$

61

where  $H_1$  and  $H_2$  show the nonlinear terms. Thus the center manifold equation has the local representation as below

$$x = h_2(y, z, \beta, \alpha_2) + O(3), \tag{6}$$

where  $h_2$  is a homogeneous polynomial of degree 2. By putting (6) in the first equation of (5), we can calculate  $h_2$  and then putting  $h_2$  in the second equation of (5), finally, after simplifying, the bifurcation is of the form

$$y' = z + A_0 \beta^2 + A_1 \beta \theta + B \beta y + C y^2, \qquad (7)$$

where

$$\begin{split} 0 &< A_0 = \left( -a\gamma_1\gamma_3 f(a,0) \frac{\partial^2 f}{\partial \beta^2}(a,0) \right) / \lambda_2, \\ 0 &> A_1 = \left( -a\gamma_3 f(a,0) \frac{\partial f}{\partial \beta}(a,0) \right) / \lambda_2, \\ B &= \gamma_1\gamma_3 \left( \frac{\partial f}{\partial \beta}(a,0) + a \frac{\partial^2 f}{\partial \beta^2}(a,0) \right) / \lambda_2, \\ C &= \left\{ (a\gamma_1\gamma_3) / (\lambda_2 f(a,0)) \right\} \left( \frac{\partial^2 f}{\partial x^2}(a,0) - \frac{\partial^2 g}{\partial x^2}(a) \right). \end{split}$$

**Proof of theorem 2.** Consider the bifurcation equation (7) with respect to the variable y=[Mdm2[t]]. The bifurcation region is determined by the sufficient condition of existence of zeroes for (7), i.e.

$$(B^2 - 4A_0C)\beta^2 - 4A_1C\theta\beta \ge 4Cz.$$

This shows that if C < 0 and  $\beta = 0$  then crossing of z = 0 to the positive area of the parameters implies a saddle node bifurcation for the system (7) and so does for (3). In this case for  $z, \beta > 0$  sufficiently small in the bifurcation area, the saddle node bifurcation causes two hyperbolic singularities of saddle type and sink type respectively. Also if z = 0 then

the bifurcation is of the form  $(B^2 - 4A_0C)\beta \ge 4A_1C\theta$  which is a straight line passing through the origin. Finally, if  $\theta = -A_0\beta/A_1$ , z = 0 then the bifurcation equation (7) is of the form  $y' = \beta y + Cy^2$  which shows a transcritical bifurcation for the system. This means that crossing the line  $\theta = -A_0\beta/A_1$  in the z=0 plane causes the transcritical bifurcation in system (3)

# **Biological results**

In the biological modeling existence of the bifurcation is important because it makes change in behavior of the system. In system (3) the level of the proteins, in particular p53, has an important role in remedy of cancer. As it is mentioned before, decline in the level of activated p53 is a certain evidence for discorrection of DNA damage. If the amount of ATM-p and the basal amount of Mdm2 are zero (i.e.  $z = \theta = 0$ ), from the discussion after theorem 1, we know that in the absence of the saddle node bifurcation (case  $v_0 < v$  in Fig. 1), the origin is the only steady state point for (3) with global attraction basin. This means that all solutions of (3) are eventually attracted by the origin (see Fig. 2).



Figure 2. Attraction of solutions to the origin for  $\nu < \nu_0$ . This occurs before the saddle node bifurcation.

This means biologically that the DNA damage remains permanently. As mentioned before, the necessary condition for damping the DNA damage signal in system (2) is that the amounts of the ATM-p protein should remains eventually nonzero. Thus, in order to repair the DNA damage, the positive amounts of the ATM-p (and also basal level of Mdm-2) must be controlled such that the system (3) passes through the saddle node bifurcation Indeed, the constants  $k_1, k_2$  in the Hill formula (see the system (2)) must be enough close to each other (this guaranties that  $0 \le |\beta|$  is small enough) and the parameter values in the system (3) must be such that  $\frac{\partial^2 f}{\partial x^2}(a, 0) > \frac{\partial^2 g}{\partial x^2}(a)$  (this guaranties that C < 0).

At the bifurcation point  $(x_1, y_1)$ , the stable manifold divides the phase space into two distinct regions (see Fig. 3). Due to the boundedness of solutions of (3), all solutions in the right hand side region are attracted by  $(x_1, y_1)$ . Similarly, as soon as the saddle node bifurcation occurs, two singularities  $(x_2, y_2)$  and  $(x_3, y_3)$  are raised for system (3) with  $x_2 < x_0 < x_3$  (see case  $v > v_0$  in Fig. 1).



**Figure 3.** Behavior at the bifurcation point  $(x_1, y_1)$  for  $v = v_0$ . The stable manifold  $W^s$  separates the phase space into two regions.

From the local bifurcation theory (see Guckenheimer, J., Holmes, P. 1993) we understand that  $x_2$  is a hyperbolic saddle point and  $x_3$  is a hyperbolic sink. In this case, the stable manifold of  $(x_2, y_2)$  divides the phase space into two distinct regions (see Fig. 4).



**Figure 4.** Behavior after the saddle node bifurcation,  $\nu > \nu_0$ . Solutions in the right hand side region are attracted by  $(x_3, y_3)$ 

All solutions in the right hand side region are attracted by  $(x_3, y_3)$ . This means that In the right hand side region of the stable manifold, the level of activated p53 remains eventually nonzero which in turn damp the DNA damage signal with a co effective work with ATM-p; whoever in the left hand side region of the stable manifold the level of activated p53 may eventually converges to zero which is a certain evidence for discorrection of the DNA damage. Thus, as it is shown in the (Fig. 3, Fig. 4), the stable manifold indicates the lower bound of amounts of p53 (and simultaneously Mdm2) for which the DNA damage signal is eventually damped.

## **Optimal Control**

Results of the previous section convince us that a fair optimal control on the system (3) must control positive amounts of the ATM-p such that the system (3) passes through the saddle node bifurcation. It must also push the solutions of the system to the attraction basin of the hyperbolic sink arisen from the saddle node bifurcation. For this purpose, we consider the positive amounts for z = [ATM - p](t) and parameters such that the system (3) passes through the saddle node bifurcation. For simplicity, we assume that  $\theta = -A_0\beta/A_1$  and consider  $\beta$  as the control function. Thus the bifurcation equation has the following form

$$y' = z + B\beta y + Cy^2.$$

The goal is finding the function  $\beta(t)$  such that  $\beta(t)$  converges approximately to zero and the solution y(t) of the bifurcation system after the sink. specific time converges to hyperbolic that is а  $y(0) = y_0, y(T) = \sqrt{-C/z}, \beta(T) \approx 0.$  Moreover  $\beta(t)$  must minimize the pass by minimizing the positive function convergence  $J = \int_{-\infty}^{T} (y - \sqrt{-C/z})^2 + \beta^2 dt$ . Thus the Hamiltonian function of the problem is of the form

$$H(y,\beta,\lambda) = \lambda(z+B\beta y+Cy^2) + (y-\sqrt{-C/z})^2 + \beta^2$$

Also the *Pontryagin's* minimum principle is  $\partial H / \partial \beta = B \lambda y + 2\beta = 0$ which implies that

$$\beta = -By \lambda / 2. \tag{8}$$

Finally the Euler-Lagrange equation of this problem is as the following form

$$y' = z + (C - \frac{1}{2}B^{2}\lambda_{2})y^{2} ,$$
  

$$\lambda_{2}' = -\lambda_{2}(2Cy - \frac{1}{2}B^{2}\lambda_{2}y) - 2(y - \sqrt{-C/z})$$
(9)

where  $\lambda(T) = 0$ . This means that the optimal function  $\beta(t)$  is found among solutions of the equations (8) and (9) simultaneously. A numerical plot for such function is shown in the Fig. 5, when the function  $\beta(t)$  is drawn for -C = B = z = 1. Obviously the solution y(t) of the bifurcation equation and the control function  $\beta(t)$  converge to  $\sqrt{-C/z} = 1$  and almost zero respectively.



Figure 5. Solutions of the optimal control

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#### === IN MEMORIAM ===

# CENTENARUL NAȘTERII PROFESORULUI DR. GEORGE EMIL PALADE (19 noiembrie 1912 – 8 octombrie 2008)

Singurul om de stiintă de origine română care a fost distins cu Premiul Nobel în Fiziologie sau Medicină, alături de Albert Claude și Christian de Duve, George Emil Palade s-a născut pe 19 noiembrie 1912, la Iași. După propriile spuse, Palade a crescut în mijlocul unei familii de profesori, învățând să îndrăgească și să respecte cărțile, să se dedice educației intelectuale. Așa cum se întâmplă adesea, tânărul George s-a orientat spre o carieră, alta decât cea sperată de tatăl său, profesor de filosofie. Palade a ales să se specializeze în Medicină, la Universitatea din Bucuresti, al cărei student a devenit în anul 1930. Ocolind conformismul și mânat de curiozitate și pasiune pentru știință, studentul medicinist Palade urmează un drum neobișnuit, concentrându-se pe aspecte fundamentale ale stiintelor medicale. Anatomia și histologia l-au interesat în mod deosebit, dedicându-se aproape în totalitate studiului microscopic al tesuturilor și celulei. Mai mult, teza sa de doctorat, pe care și-a susținut-o în 1940, a abordat un subiect iesit din tiparele preocupărilor obișnuite ale colegilor de la Medicină: structura nefronului de la delfin (Delphinus delphi) în contextul adaptării mamiferelor la viata marină. După ce a servit, în calitate de medic, Armata Română, pe durata celui de-al Doilea Război Mondial, doctorul Palade pleacă în S.U.A., în 1946. Încurajat de endocrinologul român Grigore T. Popa, descoperitorul sistemului port hipotalamo-hipofizar (Popa și Fielding, 1933), Palade a decis să emigreze și să-și continue studiile postdoctorale la Universitatea din New York, apoi la Institutul Rockefeller de Cercetări Medicale (în prezent,
Universitatea Rockefeller). Despre acest moment decisiv al carierei sale, Palade spunea într-un interviu acordat publicatiei Ad Astra în anul 2002: "Aveam sentimentul că nu știu destul, și că dacă vreau cu adevărat să fac ceva semnificativ trebuie să merg în altă parte – adică în acele locuri unde se desfășoară activități științifice cu adevărat interesante." [Tudor I. Oprea: Interviu cu George Emil Palade, Ad Astra 1(2), 2002]. La New York, Palade a fost atras de posibilitatea colaborării cu profesorul Albert Claude, un cercetător de origine belgiană, faimos deja în toată lumea pentru remarcabilele sale descoperiri biologice făcute cu ajutorul microscopiei electronice. După reîntoarcerea lui Albert Claude în Belgia, perfecționistul cercetător Palade își continuă munca neobosită de optimizare a tehnicilor de microscopie electronică, încercând să obțină preparate microscopice care să permită descrierea cât mai detaliată a organizării tisulare și celulare. Împreună cu colegii săi din "grupul Rockefeller", George Hogeboom și Walter Schneider, Palade a fost primul biolog care a apelat la fractionarea celulară ca etapă esentială în pregătirea probelor biologice. procedură ce constă în separarea organitelor celulare prin centrifugarea diferențiată a lizatului celular (Hogeboom și colab., 1948). Excelent practician, metodic și inspirat, Palade inovează în laborator, introducând tehnica păstrării intacte a organitelor celulare prin suspendarea lor în solutie izotonică de zaharoză si îmbunătățind procedeul de fixare folosind soluție de tetraoxid de osmiu tamponată la pH fiziologic (Palade, 1952). George Palade va rămâne la Institutul Rockefeller până în 1973, după care va fi invitat sa preia poziția de profesor la Facultatea de Medicină a Universității din Yale. La Institutul Rockefeller, Palade a lăsat în urmă o puternică școală de biologie celulară și moleculară, unul dintre foștii săi discipoli, Günter Blobel, fiind și el laureat al Premiului Nobel în Fiziologie sau Medicină (1999) pentru descoperirea secvențelor-semnal de transport și localizare a proteinelor (Blobel și Dobberstein, 1975a; 1975b). După 17 ani petrecuți la Yale, profesorul de biologie celulară George Emil Palade se va muta la Universitatea California din San Diego, unde a ținut cursuri în calitate de profesor emerit, până aproape de momentul decesului (7 octombrie 2008).

Considerat de către multi, drept unul dintre fondatorii "Biologiei Celulare și moleculare", Palade are meritul de a fi identificat și caracterizat, în urma unui efort consecvent, a colaborărilor cu importante nume din domeniile citologiei, histologiei, anatomiei, patologiei celulare, microscopiei si biochimiei. dar și printr-o susținere necondiționată din partea Institutului Rockefeller, o pleiadă întreagă de de structuri subcelulare. Dintre aceste descoperiri se detasează cea a particulelor ribonucleoproteice responsabile de biosinteza proteinelor. particule denumite ulterior ribosomi (corpusculii lui Palade), organite abundente în mod special în celule embrionare și glandulare (Palade, 1955). În afara acestei descoperiri majore, Palade reușește, alături de alți colaboratori, să descrie ultrastructura mitocondriei (Palade, 1953), a veziculelor sinaptice (Palade și Palay, 1954), să demonstreze natura artificială a microsomilor (Palade și Siekevitz, 1956), să caracterizeze procesele de secreție proteică din celulele pancreatice (Siekevitz si Palade, 1958a; 1958b; 1958c; 1959; 1960a; 1960b; 1962), să clarifice structura și rolurile jonctiunilor celulare de la nivelul epiteliilor (Farquhar și Palade, 1963), să descrie corpusculii Weibel-Palade cu rol în secreția factorului de coagulare von Willebrand și a selectinei P din endoteliile vasculare (Weibel și Palade, 1964), să investigheze biogeneza membranelor tilacoidale (Ohad și colab., 1967a; 1967b) sau să deslușească organizarea reticulului endoplasmatic (Leskes și colab., 1971a; 1971b), ca să amintim doar o parte din tabloul impresionantelor sale contribuții științifice.

În mod suprinzător, Palade revine în România în 1968, cu prilejul unei vizite oficiale în calitate de membru al delegației Academiei de Științe a S.U.A. Deși curtat cu intenții vădit propagandistice de către conducerea comunistă a tării, George Emil Palade pune bazele unei reale colaborări cu biologi români. În 1970, Prof. Nicolae Simionescu și Maya Simionescu au fost invitați să lucreze în Laboratorul de Biologie Celulară condus de Palade la Universitatea Rockefeller. Acest parteneriat științific, care a durat aproape două decenii, se va concretiza printr-o serie de studii ultrastructurale și fiziopatologice asupra endoteliilor capilare, dar și cu înființarea, în 1979, a Institutului de Biologie și Patologie Celulară, la București.

În aceeasi perioadă de deschidere politică și culturală a României socialiste, profesorul Palade a vizitat și Alma Mater Napocensis. În 1969. a tinut o conferință în amfiteatrul Institutului de Chimie, filială a Academiei Române, iar în luna noiembrie a anului 1971 a revenit la invitația Filialei Cluj a Academiei R.S.R. Această din urmă ocazie a prilejuit, în cadrul unei întruniri organizate la Centrul de Cercetări Biologice (C.C.B.), întâlnirea profesorului Palade cu Acad. Emil Pop, Prof. Victor Preda (directorul C.C.B. la acea vreme), Prof. V. Gh. Radu, Prof. Ioan Chiricută, Prof. Crișan Mircioiu și alții, cel mai apropiat de domeniul de cercetare fiind Dr. Constantin Crăciun, în prezent profesor, director al Centrului de Microscopie Electronică din cadrul Universității "Babeș-Bolyai". Dr. Constantin Crăciun, pe atunci având gradul de cercetător științific în Laboratorul de Biologie Experimentală al C.C.B. Clui, i-a prezentat rezultatele primelor sale cercetări ultrastructurale privind evoluția aparatului Golgi în timpul procesului de secretie muco-proteică din celulele glandulare ale canalului deferent la izopode, cercetare inspirată din lucrările profesorului Palade. Plăcut surprins, George Emil Palade, în interviul acordat prof. Mircioiu (ziarul Faclia din 4.XI.1971) a declarat că a fost "impresionat de calitatea microscopiei electronice a dr. Crăciun" și a comunicat intenția de a-l sprijini în vederea acordării unei burse în laboratorul de microscopie electronică de la Institutul Rockefeller din New York.

În continuare, Prof. Constantin Crăciun rememorează o întâmplare savuroasă petrecută cu ocazia acelei vizite, dar și alte amintiri legate de personalitatea profesorului Palade:

"A doua zi [după întâlnirea de la C.C.B., *n.a.*], profesorul Palade avea de susținut două conferințe în aula Institutului de Chimie al Academiei, ocazie cu care m-a rugat să îl ajut cu proiectarea diapozitivelor conferinței, dar, surpriză, diapozitivele sale erau pe plăci de sticlă de 9 x 12 cm, ceea ce m-a pus la grea încercare. Am reușit să găsesc în pod la Facultatea de Geografie un aparat de proiecție, dar fără rame de pus diapozitive, însă, cu ajutorul tâmplarului de aici, s-au confecționat ramele respective. După conferințe, profesorul Palade, văzând ramele din lemn și povestindu-i întâmplarea, a exclamat: "*Doamne, mintea românului!*". La întrunirea de la C.C.B., domnul Palade ne-a mai declarat: "Eu i-am extras pe soții Simionescu din România și acum lucrăm împreună". Ulterior, i-am trimis felicitări cu ocazia primirii Premiului Nobel. Prin anul 2000, i-am trimis postal câte un exemplar din revista noastră de biologie celulară [Analele Societății Naționale de Biologie Celulară, n.a.]. În iunie 2003, i-am scris personal cu referire la acordarea domniei sale, de către U.B.B., a distinctiei de "Doctor Honoris Causa", împreună cu Günter Blobel si Bhanu P. Jena, la care mi-a răspuns cu o scrisoare de multumire. În noiembrie 2003, am fost invitat împreună cu Prof. Gheorghe Benga, cu ocazia celebrării împlinirii vârstei de 90 ani a profesorului Palade, la Wayne State University, din Detroit, prilej cu care s-a instituit premiul internațional dotat cu medalia de aur "George Emil Palade", iar universitatea respectivă din Detroit a luat numele de "George Palade University". Atunci, prima medalie de aur i-a fost atribuită lui Günter Blobel, discipol de-al lui Palade, el însuși laureat al Premiului Nobel pentru medicină în 1999. În decembrie 2008, cu ocazia primirii de către mine a distincției "Doctor Honoris Causa" acordată de U.M.F. Cluj-Napoca, am închinat expunerea mea Prof. George Emil Palade, iar în octombrie 2011, la "The First World Congress on Water Channel Proteins" am prezentat expunerea "Impressions from attending in November 2003 in Detroit, U.S.A. the first "George Emil Palade" lecture and award of the "George Emil Palade" Gold Medal to Günter Blobel (1999 Nobel Laureate in Physiology or Medicine)". În anul 2011, a apărut cartea Mica enciclopedie de mari valori ridicate dintre români, în care, împreună cu Prof. Corneliu Tarba, am scris medalionul despre viata și activitatea savantului George Emil Palade. În luna martie 2013 va apărea, la editura Risoprint Cluj-Napoca, o ediție îmbunătățită a acestui volum, în limba engleză, The Encyclopedia of Outstanding Romanian Personalities, care va fi lansată la Târgul Inventică pe 19 martie, la Expo Transilvania. Pentru tot ce am scris mai sus am documente scrise însoțite de fotografii personale, inclusiv fotografia făcută de mine în 1971 profesorului Palade în timpul conferinței de la Chimie" (vezi imaginea de mai jos).



*Imagine:* Profesorul G.E. Palade conferențiind la Institutul de Chimie al Academiei Române din Cluj-Napoca despre căile de secreție ale proteinelor în celulele pancreatice (noiembrie, 1971)

(Sursa: Arhiva de Fotografii a Centrului de Microscopie Electronică, Universitatea "Babeș-Bolyai" din Cluj-Napoca).

Pe lângă Premiul Nobel obținut în 1974, Palade a mai fost onorat cu o serie de distincții, dintre care amintim premiul Fundației Albert Lasker (1966). A primit titlul de membru al Academiei de Științe a S.U.A., în 1961, iar în 1986 i s-a acordat Medalia Națională pentru Știință de către președintele Ronald Reagan. În 1976, a fost numit președinte al Asociației Americane de Biologie Celulară (The American Society for Cel Biology - ASCB), postură onorantă de care s-au mai bucurat, printre alții: Marc Kirschner (1991), J. Michael Bishop (1996), Harvey Lodish (2004) sau Bruce Alberts (2007). Pe plan național, meritele sale au fost recunoscute prin dobândirea calității de membru al Academiei Române (1975) și prin decorarea, în 2007, cu Ordinul Național Steaua României în grad de Colan, cea mai importantă distincție a statului român.

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#### === BOOK REVIEW ===

# ECOLOGICAL AND BEHAVIORAL METHODS FOR THE STUDY OF BATS, 2<sup>ND</sup> EDITION

## ENDRE JAKAB<sup>1, 2</sup>

Ecological and Behavioral Methods for the Study of Bats, 2<sup>nd</sup> edition. Edited by Thomas H. Kunz and Stuart Parsons, Johns Hopkins University Press, Baltimore, 2009, xviii + 901 p. ISBN: 978-0-8018-9147-2

Appeared after twenty-one years from the first edition the *second edition* of the *Ecological and Behavioral Methods for the Study of Bats* could be considered a primary reference for bat biologists. It is a complete guide not only for beginners but also for professional bat researchers. Edited by professor Thomas H. Kunz from Boston University, USA and bioacoustics specialist Stuart Parsons, associate professor at University of Auckland, New Zealand, this book contains comprehensive reviews written by 84 contributors from all over the world.



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### BOOK REVIEW

It is organized into eleven parts, each of them containing several chapters. Parts I to III cover methods for capturing, handling and marking bats, using conventional field techniques (such as mist netting, harp traps and use of banding devices), new radiotelemetry techniques and non-invasive ultrasonic detection methods, as well as methods for assessing colony size, prenatal and postnatal growth and development in bats, and methods for age estimation and senescence of bats.

Parts IV and V describe useful methods for studying bats in captivity as well as techniques for analysing bat morphology and migration. The following parts cover the effects of environmental contaminants, techniques for measuring milk composition in small bats (part VI), food habits analysis and nutritional ecology aspects (part VII) as well as thermal biology of bats (part VIII).

The last chapters of the book are focused on phylogenetic tools for examining evolution of bats, phylogeographic analysis of bats (part IX), methods for assessing diseases in bats and public health concerns (part X), as well as methods that promote bat conservation and a list of bat themed educational resources (part XI).

Finally I would like to thank to Carolyn Birch from Oxford Publicity Partnership Ltd. and to The Johns Hopkins University Press providing me the inspection copy of this book.