

Ultra-weak delayed luminescence in coffee seeds (*Coffea arabica* and *C. canephora*) and their germination potential: some indications for a photonic approach in seed viability

Cristiano de Mello Gallep, Evandro Conforti, members SBMO/IEEE ,
Masako Toma Braghini, Miriam Perez Maluf,
Yu Yan and Fritz-Albert Popp

Abstract— The delayed luminescence of six groups of coffee seeds were measured and analyzed by the decay behavior and statistics parameters. All groups exhibited high germination rate with normal aging behavior with exception of an unripe group. Possible correlations with the photon counting data are discussed, since all measurements exhibited higher total counting ($\Sigma(x) > 1.10^4$) indicating non-classical light storage and emission. The best correlation was observed within the total counting over the entire period of 500 points (25s), analyzed together with the decay behavior quantified by the hyperbolic decay fitting and statistical parameters.

Index Terms— Biophoton, coffee germination, delayed luminescence, photonic and life sciences

I. INTRODUCTION

THE Coffee seed normally presents high germination potential, just after appropriate harvest and desiccation. However, it loses its physiological quality very rapidly under usual storing conditions. Therefore, it is not possible to have feasible seeds, i.e. able to germinate, for more than some months. Many research groups are working for techniques to improve the seed's viability, based on data of storing conditions [1,2], methods of controlled re-hydration [3] and also low-temperature induced hibernation [4]. Although some progress has been achieved, the usual way for checking the seeds' viability and vigor is to allow them to germinate, losing so the hibernation condition. In order to distinguish between feasible and not feasible seeds, enabling an optimization of seed's storage conditions, a quick and non-destructive method is demanded, as well for other types of sensitive seeds.

In this article, an initial investigation for a biophotonic approach in the coffee seed case is presented. The delayed luminescence (DL) of seeds separated in six different groups, were measured using a special photon-counting apparatus [5].

C. M. Gallep and E. Conforti are with *Dep. Microondas e Óptica - FEEC-Unicamp*, av. Albert Einstein 400, C.P.6101, 13084-970, Campinas / SP – Brazil (55-19-37883796, gallep@dmo.fee.unicamp.br). M. T. Braghini is with *Instituto Agronômico de Campinas – Centro do Café*, C.P. 28, 13001-970, Campinas / SP – Brazil. M. P. Maluf is with *EMBRAPA - Café*, Av. W/3 Norte, sl. 322, 70770-901 Brasília, DF

Y. Yan and F.-A. Popp are with *International Institute of Biophysics*, Station Hombroich, Kapellener Strasse, D-41472 Neuss, Germany.

Thereafter, the seeds were induced to germinate under usual conditions and the germination rate was established after 15 and 30 days. The experimental data was analyzed on basis of DL-fitting parameters, statistics analysis and photon-counting distribution. The possible correlations between photon-counting behavior and seed viability are discussed, with indication for further investigation.

A. The delayed luminescence: photonic approach of vitality

The biophotonic phenomena, i.e. the ultra-weak (10-1000 photons/cm².s) delayed luminescence and spontaneous emission found in living organisms, has been studied by many multi-disciplinary groups all over the world, in a broad variety of themes. (For a good review in experimental and theoretical work, see ref. [6,7] and also the recent edition of the *Indian Journal of Experimental Biology* - May 2003, R. Bajpai ed.)

This peculiar luminescence holds much longer than the usual bio-fluorescence, and is found far from normal thermal emission, covering the entire visible spectrum and the near IR and UV. This phenomenon is still not well understood, and many arguments are displayed trying to correlate its sources, reservoirs and range of activity. Many groups, joined within the *International Institute of Biophysics* – IIB [8], believe in the holistic properties of such a biophotonic field, integrating tissues through the whole organism [9]. Recent works propose that this hypothetical macroscopic quantum channels could be based on coherent states and even squeezed states inside living organisms [10,11].

Even in research groups where the quantum-macroscopic implications of the phenomena are neglected, some correlations between the photon counting data and the subject's physiological conditions are accepted [12,13].

Recently, possible correlation between the DL behavior and the germination capacity in barley seeds was presented [14] (patent PCT/CH00/00180 [15]). Other reports are found also in ref. [6,7,16]. In the present work, a similar approach is used for coffee seeds.

II. MATERIAL AND METHODS

The DL of six groups of coffee seeds, given by the *Instituto Agronômico de Campinas* (IAC – Faz. Sta. Elisa), were measured after a five-second exposure to white-light (halogen lamp). All facilities were supported by the IIB [8], during the *Summer*

School on Biophysics, August-2003. The experimental apparatus, constituted by an automatically controlled dark chamber with a specially cooled photon-counting set-up, is well described in ref.[5]. The seeds (*Coffea arabica*, designated here *Arabica*, and *Coffea canephora*, designated here *Robusta* (commercial identifications)) were grouped, named and measured in subgroups, as presented in Table I in reverse chronological order. The size of each sub-group was determined by the seed’s size, since a constant volume was filled each time (an aluminum cuvette with front quartz window, 2 x 3.5 x 0.8 cm), in each measurement. The freshest groups (g1-g3) were also measured within their original papyrus cover protection, and the comparison with the nude measurements is discussed at the end of this work.

TABLE I - SEED’S GROUPS

Group	Harvest date	Type	Cultivar	Measur. /group	Seeds/su b-group
g1	Aug/20/03	<i>C. arabica</i>	IAC 4721	9 (14*)	15 (10*)
g2	Aug/20/03	<i>C. canephora</i>	IAC 133/c327/1653-4	9 (11*)	18 (15*)
g3	Aug/08/03	<i>C. canephora</i>	IAC 133/c298/1647-2	5 (11*)	21 (10*)
g4	Jul/15/03	<i>C. canephora</i>	IAC 133/c377/1650-6	4	25
g5	May/03/03	<i>C. arabica</i>	IAC 99	12	25
g6	Apr/20/03	<i>C. arabica</i>	IAC 144	9	20

* seeds still with natural papyrus cover

The measurements were performed under controlled temperature (22°C). In each measurement, the cuvette was filled randomly with seeds of a specific group and put inside the dark chamber. A computer software was used to control the shutters and the electronics involved in the measurement procedure, with the excitation light exposure time set to 5s, and the photon-counting started immediately after (30 ms for shutter opening/closing, light off and electronic starting), doing photon counting for the next twenty-five seconds, within 0.05-second discrete integration interval, for a total of 500 counting points. After the automatic turning-off, the cuvette was removed out of the chamber, and the procedure was repeated for all seeds. The long term DL, after more than five minutes in the dark, was also observed for each group.

In order to distinguish numerically the different decaying curves, three different approaches were used for the entire 25s period (500 points x 0.05s): 1) main statistic parameters (total number of photon counting - $\Sigma(x)$, mean, median, variance and kurtosis); 2) the best fitting parameters for a curve based on a hyperbolic-like decay, formulated by:

$$\frac{a}{(1+b)^{\alpha}} + y_0 \quad (1)$$

where $a \gg y_0$, since the photon counting is initiated just after turned off the excitation light (“a” is related to the initial values as “ y_0 ” is related to the final ones; “b” rules the short term decay velocity (\sim until $t=1$), and “ α ” rules mainly from $1 < t < 10$); and finally, 3) the counting incidence distribution, performed in order to give a graphical tool to the decaying behavior investigation.

After the dark chamber measurements, the seeds returned to Brazil and were induced to germinate under usual conditions, in the IAC facilities. In the 15th and 30th days, the germination rates were observed and registered. The germination rates, for short

(15th day) and long-term (30th day) periods, are presented in Figure 1, for the three *Arabica* and the three *Robusta* groups.

The *Arabica* groups (Fig.1a) present, in both initial and final germination rates, the aging regular behavior, as well the last two *Robusta* groups (Fig.1b). But the freshest *Robusta* group (g2, Fig.1b) presented a very low initial rate, probably due to its non-complete ripening condition.

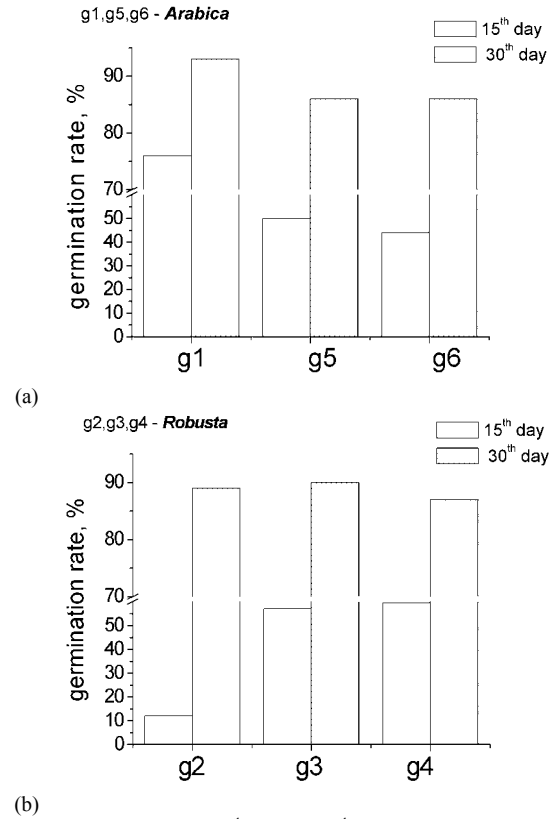


Fig.1 – germination rate for the 15th and the 30th days, *arabica* (a) and *robusta* (b) groups.

III. RESULTS AND DISCUSSION

A – Dark-counts and background

In order to compare the results and attest the set-up stability and sensitivity, the DL for the empty chamber (background) and the cuvette itself were also measured, between each seed’s group measurements. The DL curves, their incidence distribution, fitting and statistic parameters are presented in Figure 2.

As shown in Fig.2, the background measurements present very low photon counting, starting little above 100 and quickly going to the final value around 1. This behavior is indicated, besides the time curves themselves, by their parameters: low initial value ($a \sim 200$) and total counts ($\Sigma(x) < 2 \cdot 10^3$), median = 1, $\alpha = 3$ and $y_0 \sim 1$. The ultra-low DL counting can be so related to light stored in the quartz windows and inside the chamber volume itself, since just noise can be found after $t = 5$ seconds (last 380 time-points). The other case also presented in Fig.2, the empty cuvette, shows the same behavior but with a gain equally distributed in time ($a \sim 10^3$, $\alpha \sim 2.8$, $y_0 \sim 2$, $\Sigma(x) \sim 6 \cdot 10^3$, median = 3), what can be related to the additional power stored in between the cuvette

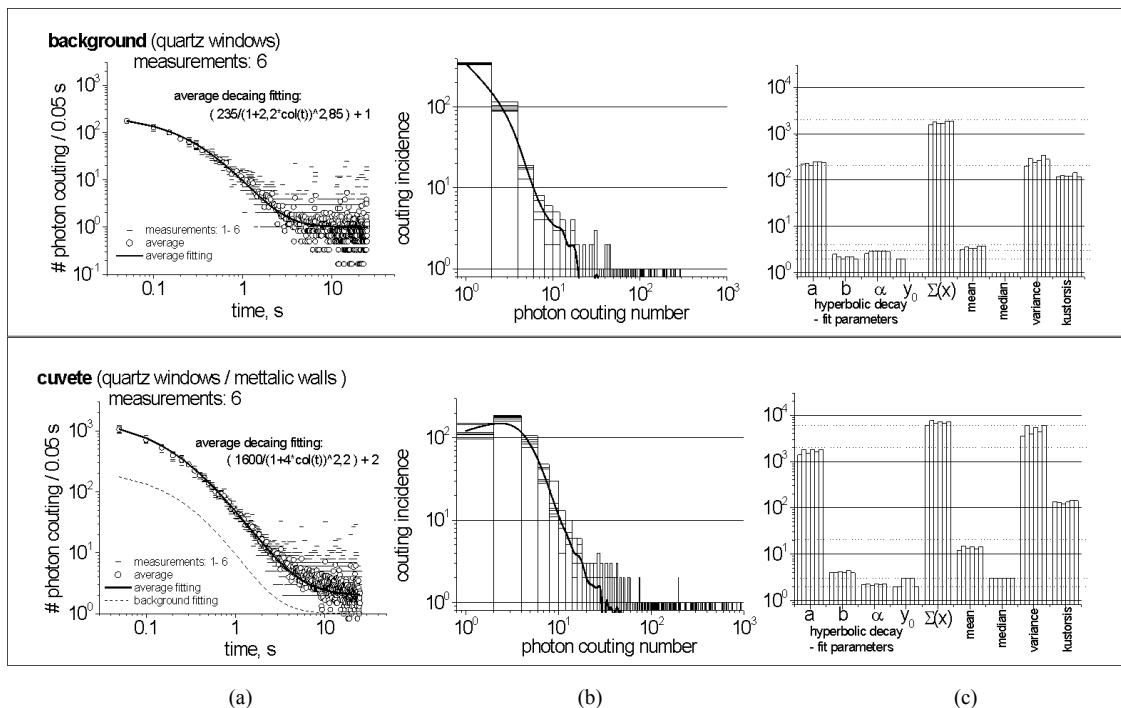


Fig.2 – DL for the background (empty chamber) and the cuvette itself: time decay and average with hyperbolic fitting (a); photon counting distribution (b); fitting and statistical parameters (c) for each measurement.

metallic walls and its quartz windows and within other chamber’s windows. The cuvette was than chosen as the equivalent background in this experimental situation.

In both these two cases, the kurtosis is high (~ 100), as they have a pronounced distribution peak, and the variance goes with the a factor, since the curve is a decay one.

B – The *Coffea arabica* and *Coffea robusta* groups

The photo-counting data for the three *Arabica* groups are presented in Figure 3, as done in Fig.2 for background and cuvette. The $g1$ data shows a pronounced difference from de cuvette data, with lower initial counting values and much more high final values, reached after a slower decay ($\alpha < 2$, $y_0 \sim 10$, median ~ 10). This behavior is nicely illustrated in the counting distribution (Fig.3b($g1$)) and by the total counting ($\Sigma(x) \sim 1.10^4$). For the other two groups, $g5$ and $g6$, the main difference from the cuvette data is noted mainly after $t = 5s$, and so they also present high $\Sigma(x)$, but with lower final value than $g1$, (median ~ 4 and y_0 between 3 and 4).

All three *Arabica* groups exhibit distinguishable data from the cuvette, mainly for $t > 5s$, with similar $\Sigma(x)$ but distinguishable medians/ y_0 . No usual short-term photoluminescence as observed, for example, in leaves and other tissues with chlorophyll, was measured [6,7,11], indicating that seeds could be in the desired dormant condition, with the predominant long-term photon-counting decay ($t > 5s$).

The photon-counting data for the three *Robusta* groups are presented in Figure 4. A new behavior is found in the $g2$ group: a very slow decay ($\alpha \sim 1.3$) was detected, as observed in experiments with leaves [6,7,11]. The $g2$ sub-groups present the highest initial values ($a \sim 2.8 \cdot 10^3$) found in this work, but tending to a low final one ($y_0 \sim 4$). They also present

the highest total count ($\Sigma(x) \sim 5.10^4$, $20 < \text{median} < 30$), and the photon counting distribution is broader, giving the smallest kurtosis (~ 6). All these data indicate that the evaluated seeds could be yet unripe and have not entered into the dormant state. The other two groups, $g3$ and $g4$, present decays similar to the *Arabica* ones, with noted high final values (for $g3$: $y_0 \sim 16$, $\Sigma(x) \sim 2.10^4$, median ~ 18 , and for $g4$: $y_0 \sim 7$, $\Sigma(x) \sim 10^4$, median ~ 9). These behavior could indicate that these groups are well ripen and with high viability. As found for the three previous groups, all *Robusta* groups also showed distinguishable data in long-time photo-counting terms ($t > 5s$).

The longer term DL, checked for all groups, holds for minutes (measured from 5 to 10 min. in dark), with very low (10%-20%) decreasing. It indicates that longer term DL could be a more efficient approach to check seed viability, at least for the coffee case. Looking back at Fig.1, simultaneously with Fig.3 and Fig.4, it is found that the $\Sigma(x)$ could be tested as a qualitative indicator for germination capacity, and the final DL value (quantified by the median and y_0) could be tested as a quantitative indicator. The exception to this simple approach, group $g2$, needs a more elaborated treatment, involving the α parameters to quantify the non-usual (for seeds) slower DL. As mentioned in the section II, the first three groups were measured also with seeds within their natural clad, formed by dead tissues and looking like a fine paper coating (*papirus*). For all the three cases (not shown here), the DL fitting tends to the same behavior as the “nude” ones after 5 seconds, differing from it by a factor lower than 2. This comparison indicates that is not necessary to remove the seed’s protection to execute this type of measurement, enabling seed testing without any damage to the seed.

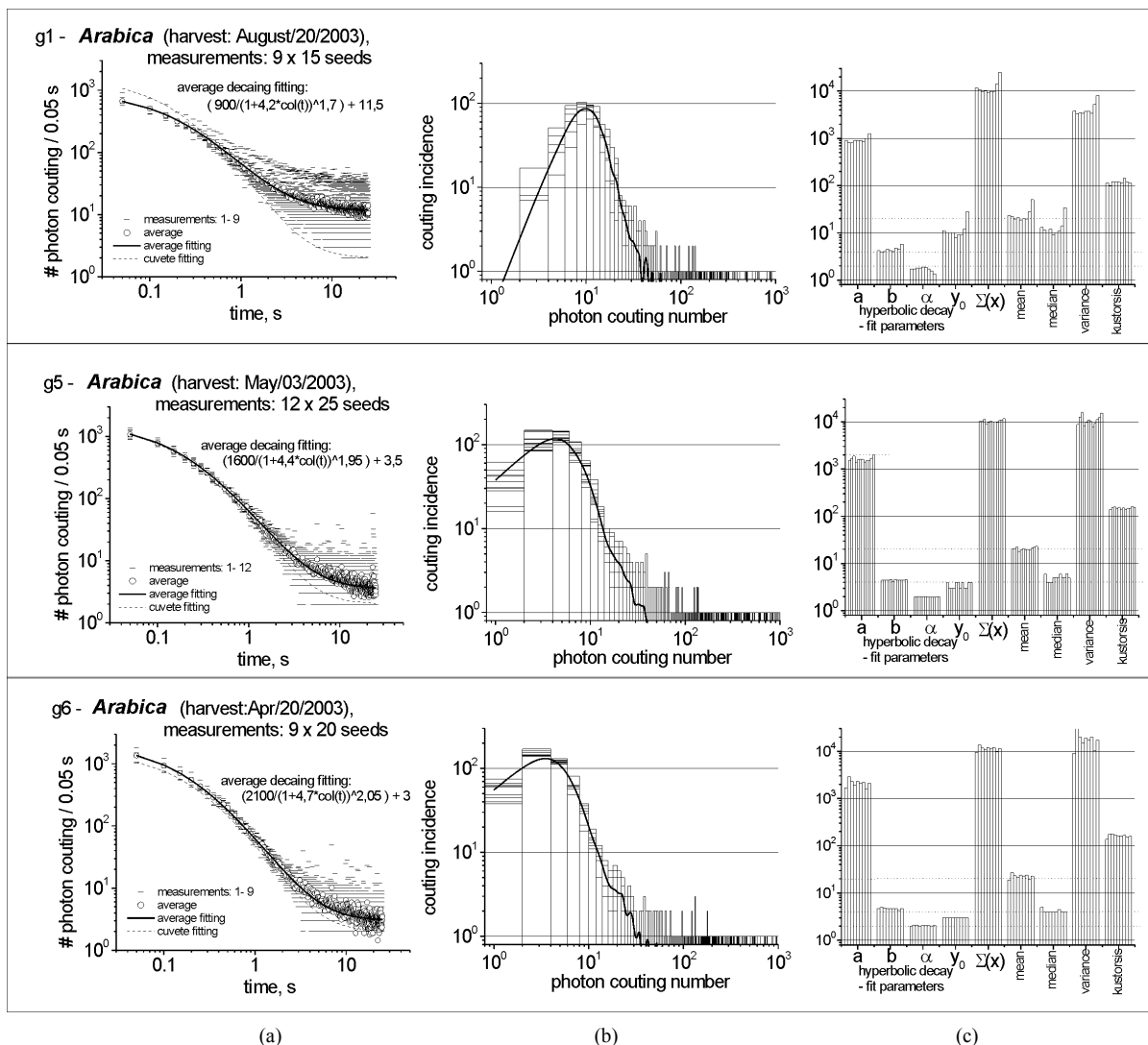


Fig.3 – DL for the three Arabica groups (*g1*, *g5* and *g6*): time decay and average with hyperbolic fitting (a); photon counting distribution (b); fitting and statistical parameters (c) for each measurement. The average cuvette fitting is also plotted in (a).

IV – CONCLUSION

The ultra-weak Delayed Luminescence measurements and analysis data of coffee seeds from three *C. arabica* and the three *C. canephora* different groups, all with high final germination capacity, presented sometimes small but always detectable differences with the empty cuvette data. All coffee DLs exhibited high total counting ($\Sigma(x) > 1.10^4$), indicating non-classical light storage and/or emission, as usually found for plants in the specific literature. A leaf-like behavior was found in one freshest group, indicating an unripe condition, which was corroborated by the group's low germination capacity in short-term regime (15 days).

The authors believe that the present work shows some parameters related to the DL in seeds, which could be more studied as possible keys in a photonic approach of viability. The best correlation with the germination capacity was found within the total counting $\Sigma(x)$ over the entire period of 500 points (25s), together with the DL decay behavior, illustrated by the DL time decay itself and by the photon counting distribution, and numerically quantified by hyperbolic decay

fitting and also by statistics parameters. Further work should consider broader groups of coffee seeds, by varying their age, harvest condition and even trying artificial stress, with possible correlated photon burst detection. The technical viability of a practical photonic method in this field needs to be carefully analyzed, since can promote unpredictable new functionalities in storing methods and related agricultural research.

ACKNOWLEDGMENT

The authors would like to send their gratitude to the International Institute of Biophysics, and to prof. Popp, Sophie Cohen and Yu Yan, for promoting the summer schools and these specific measurements. C. M. Gallep would like also to acknowledge all people, met in the occasions of the IIB summer schools, for their informal contributions to this work. Special thanks to Manish Vekaria and the students from the Korean group. The participation in the third IIB Summer School was partially supported by FAEP/Unicamp.

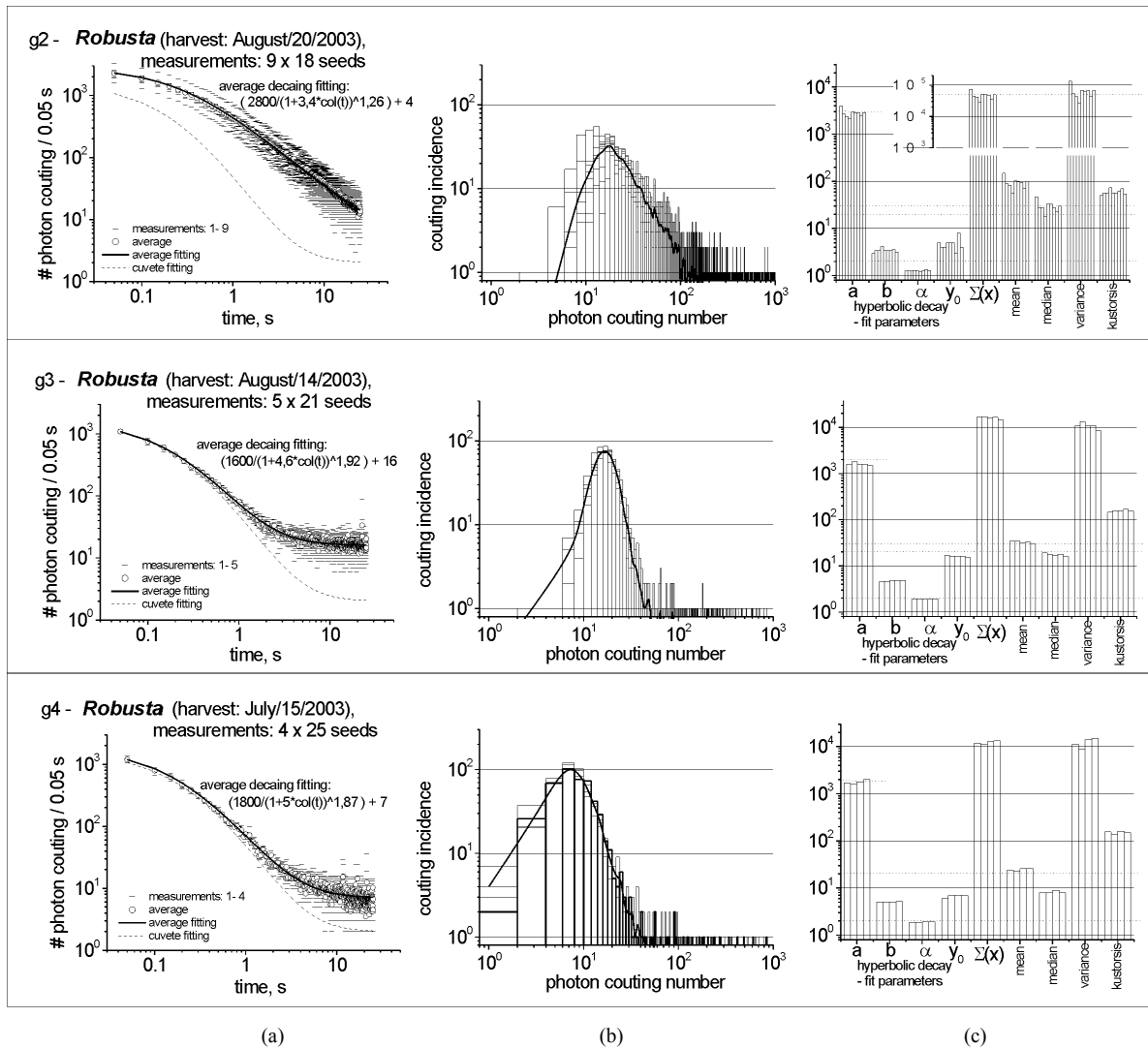


Fig.4 – DL for the three Robusta groups (g2, g3 and g4): time decay and average with hyperbolic fitting (a); photon counting distribution (b); fitting and statistical parameters (c) for each measurement(c). The average cuvette fitting is also plotted in (a).

REFERENCES

- 1 - E. Couturon, “Maintaining the viability of coffee seeds by checking their water-content and storage-temperature”, *The Cafe Cacao*, Vol.24 (1980), No. 1, pp. 227-32.
- 2 - T. D. Hong, R.H. Ellis, “Optimum air-dry seed storage environments for arabica coffee”, *Seed Science and Techn.*, Vol. 20(1992), No. 3, pp. 547-560.
- 3 - S. Dussert *et al.*, “Beneficial effect of post-having osmoconditioning on the recovery of cryopreserved coffee () seeds”, *Cryo-Letters*, Vol.21 (2000), No. 1, pp. 47-52.
- 4 - S. Dussert *et al.*, “Tolerance of coffee () seeds to ultra-low temperature exposure in relation to calorimetric properties of tissue, water, lipid composition and cooling procedure”, *Physiologia Plantarum*, Vol.112(2001), No. 4, pp. 495-504.
- 5 - F.A. Popp, “Biophotons – Background, experimental results, theoretical approach and applications”, *Res. Adv. Photochem. & Photobiol.*, Vol.1(2000), pp. 31-41.
- 6 - J.J. Chang, J. Fisch e F.A. Popp (ed.), *Biophotons*, Kluwer Acad. Publ., 1998.
- 7 - multi-author review editions of *Experientia*, Vol.44 (1988), pp. 543-600, and Vol.48 (1992), pp. 1029-1102.
- 8 - Neuss – Germany: www.lifescientists.de
- 9 - R. Wijk *et al.*, “Regulatory aspects of low intensity photon emission”, *Experientia*, Vol.44 (1988), pp. 586-593.
- 10 - F.A. Popp e Y.Yan, “Delayed Luminescence of Biological Systems in Terms of Coherent States”; and F.A. Popp, J.J. Chang, A. Herzog, Z. Yan e Y. Yan, “Evidence of Non-Classical (Squeezed) Light in Biological Systems”, *Physics Letters A*, Vol. 293 (2002), pp.93-97 and pp.98-102 respectively.
- 11 - R. P. Bajpai, “Coherent nature of the radiation emitted in delayed luminescence of leaves”, *J. Theor. Biology*, Vol.198 (1999), pp.287-299.
- 12 - T. Ohia *et al.*, “Biophoton emission due to drought injury in red beans: possibility of early detection of drought injury”, *Jpn. J. Appl. Physics*, Vol. 41(2002), No.7A, pp. 4766-4771.
- 13 - M. Kobayashi e H. Inaba, “Photon statistics and correlation analysis of ultraweak light originating from living organisms for extraction of biological information”, *Applied Optics*, Vol.32 (2000), No.1, pg. 183-192.
- 14 - Y. Yan., F.A. Popp, G.M. Rothe, “Correlation between germination capacity and biophoton emission of barley seeds (*Hordeum vulgare L.*)”, *Seed Science and Techn.*, Vol. 31(2003), No. 2, pg. 249-258
- 15 - F.A. Popp, “Method, system and use of measuring devices for determining the germinability of seeds” PCT/CH00/00180, European Patent Office (EP1 188 041 B1).
- 16 - J.M. Souren, E.K.Boon-Niermiejer, R. Van Wijk, “Photon emission and germination capacity of tomato seeds”, *Conference on Biophotons 1999*, Neuss/Germany, in www.lifescientists.de/ib_004e38.html.