Brazilian cerrado antioxidant sources: cytotoxicity and phototoxicity in vitro

Fontes de antioxidantes do cerrado brasileiro: citotoxicidade e fototoxicidade in vitro

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Abstract

Annona crassiflora (araticum), Eugenia dysenterica (cagaita), and Caryocar brasiliense (pequi) are tropical fruits of the second biggest Brazilian biome: the cerrado. Nowadays, the cerrado faces two different realities: 1) the great possibility of food production since it is considered as the biggest storehouse of the world; and 2) the rich biodiversity that has been newly discovered and known. Previous studies showed that certain cerrado fruits demonstrate high content of total phenols and excellent antioxidant activity in in vitro models. Moreover, using fingerprinting analysis, important bioactive molecules were identified as probably responsible for their antioxidant activity. In this study, the cytotoxicity and phototoxicity of ethanolic extracts from cerrado fruits were evaluated using the in vitro Neutral Red Uptake (NRU). Regarding cytotoxicity, the extracts of araticum peel and cagaita seed did not show any cytotoxic potential up to 300 µg.mL⁻¹. Ethanolic extracts of araticum seed and pequi peel presented low cytotoxic potential and, according to linear regressions, the estimated LD₅₀ were 831.6 and 2840.7 mg.kg⁻¹, respectively. In the evaluated conditions, only the araticum peel extract presented a phototoxic potential. This is the first attempt to screen the toxicity of cerrado fruits with high antioxidant activity.

Keywords: tropical fruits; cerrado; antioxidant activity; 3T3 neutral red uptake; cytotoxicity; phototoxicity.

1 Introduction

The importance of the search for and exploitation of natural antioxidants, especially of plant origin, has greatly increased in recent years (JAYAPRAKASHA; RAO, 2000). Both epidemiological and clinical studies have provided evidence that phenolic antioxidants present in cereals, fruits, and vegetables are the principal contributing factors in accounting for the significant reduced incidences of chronic and degenerative diseases encountered by populations whose diet is high in the intake of these foods (SHAHIDI, 1996). Typical compounds that possess antioxidant activity include phenols, phenolic acids and their derivatives, flavonoids, tocols, phospholipids, amino acids and peptides, phytic acid, ascorbic acid, pigments, and sterols (SHAHIDI, 1996). A phenol contains an –OH group attached to a benzene ring. Many phenols exert powerful antioxidant effects in vitro, inhibiting lipid peroxidation by acting as chain breaking peroxyl radicals’ scavengers. Phenols with two adjacent –OH groups, or other chelating structures, can also bind transition metal ions in forms poorly active in promoting free-radical reacting. Phenols can also directly scavenge Reactive Oxygen Species (ROS), such as OH, ONOO⁻, and hOCl (HALLIWELL; GUITTERIDGE, 1999).

Annona crassiflora (araticum), Eugenia dysenterica (cagaita), and Caryocar brasiliense (pequi) are tropical fruits of the second biggest biome of Brazil: the cerrado. Although these fruits are highly used by traditional herbal medicine such as herbal healing, there are few scientific studies on the relationship between the biological activities and the ethnobotanical uses in order to validate them. Our previous studies show that ethanolic extracts of these fruits demonstrated high content...
of total phenols and excellent antioxidant activity using the
in vitro model 2,2-diphenyl-1-picrylhydrazyl (DPPH) and
chemically induced lipid peroxidation using rat liver microsome
as oxidative substrate. The best results were found for ethanolic
extracts of pequi peel, cagaita seed, araticum seed, and araticum
peel. The IC$_{50}$ results were in the range of 14 to 50 µg.mL$^{-1}$
for DPPH assay and 0.7 to 4.5 µg.mL$^{-1}$ in the lipid peroxidation
model. The use of direct infusion electrospray ionization mass
(ESI-MS) and tandem Mass Spectrometries (MS) revealed the
presence of important bioactive molecules in the cerrado fruit
extracts, probably responsible for their antioxidant activity,
such as ascorbic acid, gallic acid, quinic acid, dicafeoylquinic
acid, quercetin, quercetin 3-O-arabinose, ferulic acid,
rutin, xanthohyolin, caffic acid, and its derivatives such as
cafeoyltartaric acid and caffeoyl glucose (ROESLER et al.,
2006, 2007a, b).

In order to ensure that all potentially hazardous products,
especially new substances, have proper warning labels and use,
regulatory agencies require determination of acute toxicity
hazard potential of substances and products. This determination
is currently made using a test that requires laboratory rats.
Historically, lethality estimated by the dose of a substance that
produces 50% of animal death (LD$_{50}$) has been the primary
toxicological endpoint in acute toxicity tests. The conventional
LD$_{50}$ acute oral toxicity in vivo test method has been modified
in various ways to reduce and refine animal use in toxicity
testing (ORGANISATION..., 2001; ENVIRONMENTAL....,
2002). In vitro cytotoxicity methods have been evaluated as
other means to reduce and refine the use of animals, and these
methods may be helpful in predicting in vivo acute toxicity.
Since moving the starting dose closer to the LD$_{50}$ reduces
the number of animals necessary for the acute oral systemic toxicity
test, the use of in vitro cytotoxicity assays to predict a starting
dose close to the LD$_{50}$ may reduce animal use. The Interagency
Coordinating Committee on the Validation of Alternative
Methods (ICCVAM) and the National Toxicology Program
(NTP) Interagency Center for the Evaluation of Alternative
Toxicological Methods (NICEATM) recommended further
evaluation of the use of in vitro cytotoxicity data as one of the
factors used to estimate the starting dose for in vivo acute lethality studies based on the preliminary information that this
approach could reduce the number of animals used in in vivo
studies (i.e., reduction), minimise the number of animals that
receive lethal doses (i.e., refinement), and avoid underestimating
hazard (NATIONAL INSTITUTE...., 2001).

Cytotoxicity has been defined as the adverse effects resulting
from the interference with structures and/or processes essential
for cell survival, proliferation, and/or function (EKWALL,
1983). Other researchers (GRISHAM; SMITH, 1984) also
concluded that since the actions of substances that produce
injury and death are ultimately exerted at cellular level in
vitro cytotoxicity assays may be useful for the prediction of
acute lethal potency. The endpoint measured in the in vitro
Neutral Red Uptake (NRU) cytotoxicity test is cell viability and
the major endpoint of interest is the half maximal inhibitory
concentration, i.e. the concentration which promotes 50% of
inhibition in the NRU (IC$_{50}$).

The dermal phototoxicity of a chemical is defined as a
/toxic response that is elicited after the exposure of skin to
a chemical or systemic administration of a chemical and
subsequent exposure to Ultraviolet (UV) light. As demonstrated
by several validation studies (BORENFREUND; PUERNER,
1985; CENTRE....; EUROPEAN...., 2000; NATIONAL INSTITUTE....,
2001), the phototoxic potential of chemicals can be effectively assessed by in vitro methods. In 1991, the
European Commission (EC), initially represented by the
Directorate General XI and later by the European Centre for the
Validation of Alternative Methods (ECVAM) and the European
Cosmetic, Toiletry and Perfumery Association (COLIPA), agreed
to establish a joint EU/COLIPA programme on the development
and validation of in vitro phototoxicity tests. In 1996, the
Organisation for Economic Co-operation and Development
(OECD) recommended an in vitro tier-testing approach for
phototoxicity assessment (SPIELMANN et al., 1998). In 2000,
the Commission of European Communities put into force the
Directive 200033EC, which introduces the in vitro 3T3 NRU
phototoxicity test as a validated replacement for testing methods
involving the use of laboratory animals. The essence of this test
is to compare the cytotoxicity of a chemical when tested in the
presence and absence of exposure to a non-cytotoxic dose of
ultraviolet A (UVA) light (PETERS; HOLZHUTTER, 2002).

The aim of this study is to evaluate the toxicity of the Brazilian
cerrado fruit extracts by using the in vitro 3T3 NRU cytotoxicity
and phototoxicity methods in order to estimate the applicability
of these fruits as a sustainable source of natural antioxidant for
cosmetic, foods, and pharmaceutical applications.

2 Materials and methods

2.1 Cell lines

BALB/C 3T3 mouse fibroblasts, clone 31, were obtained
from ATCC (Manassas, VA, USA).

2.2 Chemicals and media

Dulbecco’s Modified Eagle Medium (DMEM) (Gibco, Grand
Island – NY, USA), Phosphate Buffered Saline (PBS), NR dye
(Sigma-Aldrich, Saint Louis, MO, USA), New Born Calf serum
(NBCS) (Nutricell, Campinas – SP, BR), acetic acid glacial
(Carlo Erba, Rodano, MI, IT), sodium laurel sulphate (positive control for cytotoxicity) (Sigma-Aldrich,
Saint Louis, MO, USA), and chlorpromazine (positive control for
phototoxicity) (Sigma-Aldrich, Saint Louis – MO, USA).

2.3 Plant material

Annona crassiflora (araticum), Eugenia dysenterica
(cagaita), and Caryocar brasiliense (pequi) were selected based
on the traditional application and lack of information related to
acute toxicity (Table 1). They were obtained from Erlow Farm,
Km 07 Br 070, Goiânia, Center West Brazil. Fruit harvests were
conducted twice and special care was taken to avoid damaging
the fruits. The fruits were transported to the State University of
Campinas – UNICAMP and stored at 5 °C until use for
two months maximum.
2.4 Ethanolic extraction

Undamaged fruits were selected and peeled, and the pulp and seeds were manually separated. About 100 g of each part was cut in small fragments, mixed, and extracted twice with 300 mL of aqueous ethanol (5:95, v/v, water: ethanol) using a household mixer for 20 minutes. The extracts were filtered through a cotton membrane and the residue was re-extracted under the same conditions. The residues were kept for further evaluations. The resultant material was subjected to vacuum rotary evaporation at 40 °C to remove ethanol. The concentrated ethanolic extracts were lyophilised and stored at −18 °C in amber glass bottles until use.

2.5 In vitro cytotoxicity

The assays were carried out according to the official protocol from OECD (ORGANISATION..., 2001). Briefly, 3T3 cells were seeded into 96-well plates to form a sub confluent monolayer (1 × 10^4 cells/well). The culture medium was removed and different concentrations of test chemicals in medium were added to cells for 48 hours, 37 °C, 5% CO₂. The untreated vehicle control was the same as the treatment medium. The cells were washed once with PBS, and NR dye medium was added for 3 hours, 37 °C, 5% CO₂. The cells were washed with PBS, and NR desorb solution (50:1:49, v/v/v, ethanol: acetic acid: water) was added to the plates. The plates were shaken for 20 minutes and NR absorption was measured at Optical Density (OD) 540 ± 10 nm. Cell viability was calculated as the percentage of control values to define IC_{50}, substance concentrations (µg.mL⁻¹) (NATIONAL TOXICOLOGY..., 2003, 2006).

2.6 In vitro phototoxicity

The assays were performed according to the official protocol from NIH (NATIONAL INSTITUTE..., 2001). Briefly, the cells were prepared as in the cytotoxicity protocol considering two plates for each test sample. The 3T3 NRU test for phototoxicity requires a 60-minute exposure to sample dilutions in PBS followed by UVA (5 J.cm⁻²) exposure of one of the plates (the other plate is used as non irradiated control). After washing both plates with PBS, the cells were incubated for 24 hours, 37 °C, 5% CO₂. The plates were washed, incubated with NR dye medium, and re-washed. The NR desorb solution was added and NR absorption was measured at OD 540 ± 10 nm. Cell viability was calculated for each treatment. Phototoxicity was assessed by comparing the differences in toxicity between negative control plates that had not been exposed to UVA and test plates exposed to UVA. The Photo-Irritancy Factor (PIF) relates the half-effective concentration values (EC_{50}) of the curve for darkness (EC_{50}[-UV]) and in the presence of light (EC_{50} [+UV]) by means of the following equation (Equation 1):

\[
PIF = \frac{EC_{50}([-UV])}{EC_{50}([+UV])}
\]

Depending on whether the PIF value is larger or smaller than the cut-off value of five units, the chemical is classified as phototoxic or non-phototoxic, respectively. A shortcoming of the measure in Equation 1 is that additional ad hoc definitions are required to cope with situations in which no EC_{50} values can be derived from the corresponding concentration-response curve: 1) if no EC_{50} value can be derived from one of the curves, the corresponding EC_{50} value in Equation 1 is replaced by the highest concentration tested, and the chemical is classified as phototoxic if this modified PIF value is larger than one unity; and 2) if no EC_{50} value exists for both curves, the chemical is considered non-phototoxic (Borenfreund; Puerner, 1985; CENTRE..., EUROPEAN..., THE EUROPEAN..., 2000; NATIONAL INSTITUTE..., 2001).

A second measurement of the difference between the dark and light curve, the so-called Mean Photo Effect (MPE), was proposed by Holzhüttter (1997). It aims to overcome the obvious limitations in the application of the PIF by comparing the two curves at arbitrary doses, and it is useful even in situations where no EC_{50} values can be derived from the corresponding concentration-response curve. The MPE measures the effect of UV exposure over a range of concentrations. A material is considered non-phototoxic if the MPE is < 0.1 and phototoxic if the MPE is ≥ 0.1.

2.7 Data analysis

Both measures of curve difference, PIF and MPE, represent statistical estimates that have to be derived from the observed concentration-response relations. To facilitate this work and to harmonise the process of data analysis, the software PHOTOTOX 2.0 (ZEBET, Diedersdorfer, BE, DE) was used. This program module performs a bootstrap resampling of the original concentration-response data that result in a set of new computer-generated concentration-response data (100 times), which can be considered as equally probable realisations of the “true” concentration-response data hidden in the experimental observations. Based on these curves, and at given cut-off values, the program calculates PIF and MPE, and determines the probability (p-value) that a test chemical is phototoxic (PETERS; HOLZHÜTTER, 2002).

Table 1. Ethnobotanical data of selected cerrado fruits.

| Specie                                | Family          | Local name          | Portion used | Traditional uses                      |
|---------------------------------------|-----------------|---------------------|--------------|---------------------------------------|
| Annona crassiflora                   | Annonaceae      | Araticum, Marolo,   | Seed         | Snake bite(2), hair antiparasitic(2)   |
| Annona rodrigueisi                   |                 | Cabeça de negro     | Seed, leaves | Anti diarrheas(2,3)                    |
| Eugenia dysenterica                  | Myrtaceae       | Cagaita             | Fruits       | Lexitive(2)                           |
| Stenocalyx sp.                       |                 |                     | Leaves       | Anti diarrheas(2)                      |
| Caryocar brasiliense                 | Caryocaraceae   | Pequi, piqui, piquia| Nut oil      | Asthma, influenza, cold, bronchopulmonary diseases(2,3) |
|                                      |                 |                     | Seed         | Aphrodisiac tonic(2)                  |
|                                      |                 |                     | Leaves       | Regulation of women period(2)         |

*These data was based on 1) Weinberg et al. (1993); 2) Almeida et al. (1998); and 3) Rodrigues e Carvalho (2001).
2.8 Prediction of LD$_{50}$

To obtain a model for the prediction of lethal dose LD$_{50}$ values from IC$_{50}$ values, Halle (2003) calculated a linear regression from pairs of the log-transformed IC$_{50}$ (in mM) and log transformed rodent oral LD$_{50}$ values (in mmol.kg$^{-1}$). Equation 2, which was obtained from a correlation between values of in vitro cytotoxicity and rat and mouse oral LD$_{50}$ values for 347 chemicals, is:

$$\text{Log LD}_{50} (\text{mmol.kg}^{-1}) = 0.435 \times \log \text{IC}_{50} (\text{mM}) + 0.625$$  \hspace{1cm} (2)

This Equation was denominated RC regression. Presumably, the substance units were expressed in moles because these are units that produce biological activity and, hence, they are expected to produce the best fitting regression. IC$_{50}$ values for 347 substances were obtained from 157 original publications (HALLE, 2003). The 1912 IC$_{50}$ values, two to 32 per substance, were averaged using geometric means to produce one IC$_{50}$ value for each substance.

To improve the RC regression in terms of the prediction of LD$_{50}$ values by in vitro NRU IC$_{50}$ values, some regressions were developed using RC data in weight units to excluding: 1) mouse data (i.e. RC rat-only regression); and 2) substances with mechanisms of toxicity that were not expected to be active in the 3T3 cell cultures (i.e. RC rat-only regression excluding substances with specific mechanisms of toxicity) (NATIONAL TOXICOLOGY..., 2006).

Equation 3 for the RC rat-only weight regression is:

$$\text{Log LD}_{50} (\text{mg.kg}^{-1}) = 0.372 \times \log \text{IC}_{50} (\text{µg.mL}^{-1}) + 2.024$$  \hspace{1cm} (3)

Equation 4 for the RC rat-only weight regression excluding substances with specific mechanisms of toxicity is:

$$\text{Log LD}_{50} (\text{mg.kg}^{-1}) = 0.357 \times \log \text{IC}_{50} (\text{µg.mL}^{-1}) + 2.194$$  \hspace{1cm} (4)

2.9 Solubility protocol

For testing the chemicals prepared in the chemical dilution medium, the highest test article concentration that may be applied to cell in the main experiments will be either 100 mg.mL$^{-1}$ or the maximum soluble dose. For test chemicals prepared in either DMSO or ethanol, the highest test article concentration that may be applied to the cells in the major experiments will be either 2.5 mg.mL$^{-1}$ or less depending upon the maximum solubility in the solvent (NATIONAL TOXICOLOGY..., 2003, 2006).

3 Results and discussion

3.1 Cytotoxicity

NR is a weakly cationic, water-soluble dye that stains living cells by readily diffusing through the plasma membrane and concentrating in lysosomes. The intensity of the dye in the culture is directly proportional to the number of living cells. In addition, since altering the cell surface or the lysosomal membrane by a toxicological agent causes lysosomal fragility and other adverse changes that gradually become irreversible, cell death and/or inhibition of cell growth decreases the amount of NRU by culture (BORENFREUND; PUERNER, 1985).

A calculation of cell viability is expressed as NRU for each concentration of the chemical by using the mean NRU of four replicate values (blank is subtracted). These values are compared with the control mean values. Relative cell viability is then expressed as a percentage of untreated vehicle control.

Cagaita seed (Figure 1a) and araticum peel (Figure 1c) ethanolic extracts did not decrease cell viability up to 300 µg.mL$^{-1}$. Thus, no significant general toxicity was observed and the IC$_{50}$ values higher than 300 µg.mL$^{-1}$. Due to the low solubility of these fruit extracts, it was not possible to increase the concentration in order to obtain an IC$_{50}$ value. Pequi peel extract was tested up to 3000 µg.mL$^{-1}$. At this concentration, the pequi peel extract started producing a precipitated in the cell culture medium and caused a decrease in cell viability. If the linear correlation of cell viability decrease in the PHOTOTOX program and pequi peel ethanolic extract concentration are considered, the mean IC$_{50}$ would be 3731.3 ± 579 µg.mL$^{-1}$ (Figure 1d). Regarding the araticum seed ethanolic extract, the results suggested some evidence of moderate toxicity, and mean IC$_{50}$ value for the araticum seed ethanolic extract was 108.0 ± 7 µg.mL$^{-1}$ (Figure 1b).

Using the regressions previously described, the LD$_{50}$ values for the extracts were predicted. Considering Equation 3 (RC rat-only weight regression), the LD$_{50}$ result obtained for the pequi peel extract was 2169.4 mg.kg$^{-1}$, and for the araticum seed extract it was 603.2 mg.kg$^{-1}$. Using Equation 4 (RC rat-only weight regression excluding substances with specific mechanisms of toxicity), the LD$_{50}$ results were 2840.7 and 831.6 mg.kg$^{-1}$ for the pequi peel and araticum seed extracts, respectively. For comparative purposes, Table 2 shows some chemicals’ data regarding LD$_{50}$ and IC$_{50}$.

Previous studies in vivo with two groups of seven rats each showed that orally administrated aqueous solution of A. crassiflora seed extract (50 mg of gallic acid equivalents per kg, i.e. 767 mg of crude extract per kg of weight) for 14 consecutive days did not produce any death, clinical signs, or gross alterations attributable to hepatic toxicity or other organ damage if compared to a control group that received saline solution until

| Extract or chemical compound | Cytotoxicity IC$_{50}$ (µg.mL$^{-1}$) | Oral LD$_{50}$ (mg.kg$^{-1}$) |
|-----------------------------|-----------------------------------|-------------------------------|
| Cagaita seed                | > 300.0                           | ND                            |
| Araticum seed               | 108.0 ± 7                         | 831.6                         |
| Araticum peel               | > 300.0                           | ND                            |
| Pequi peel                  | 3731.3 ± 579                      | 2840.7                        |
| Sodium lauryl sulphate      | 40 ± 0$^{(2)}$                    | ND                            |
| Phenol                      | 50 ± 10$^{(2)}$                   | 414$^{(2)}$                  |
| Nicotine                    | 270 ± 70$^{(2)}$                  | 50$^{(3)}$                    |
| Glycerol                    | 24650$^{(2)}$                     | 12691$^{(3)}$                 |
| Bergamot oil                | ND                                | ND                            |
| Caffeine                    | ND                                | 192$^{(2)}$                   |

The oral LD$_{50}$ values were calculated for the plant extracts using the RC rat-only weight regression. The different compounds in the table are references for cytotoxic potential comparison. These data were based on 1) National Toxicology... (2006); 2) Ghs (GLOBALLY..., 2005); and 3) Halle (2003). ND: not determined.
Cerrado antioxidants: toxicity in vitro

Similar since animals are comprised of organ systems consisting of tissues, which are comprised of cells. All cells, regardless of whether they are animal cells or in vitro cell cultures, have similar cellular mechanisms of energy production, utilisation, and maintenance of cell membrane integrity. Animal death and death of cells in culture due to toxicity are similar in that both involve some type of injury. For the animals, the cellular injury leads to tissue and organ injury to the most sensitive target organ, which may then cause the death of the whole organism. An organ system failure can be due either to the death of cells in the affected organ or to the loss of function of the surviving cells in the organ, which results in cell death or loss of function in other organs (GENNARI et al., 2004). The death of an animal is a result of major organ system failure. Ultimately, the cardiovascular and respiratory systems fail. Respiratory depression may be due to the depression of the central nervous system rather than a direct assault on the respiratory system. Other major organ system failures, such as liver and kidney failure, gastrointestinal corrosion, and bone marrow depression, also cause death. Cell death is a culture system that involves the death of a single cell type. Cell death and animal death may be similar since animals are comprised of organ systems consisting of tissues, which are comprised of cells. All cells, regardless of whether they are animal cells or in vitro cell cultures, have similar cellular mechanisms of energy production, utilisation, and maintenance of cell membrane integrity. Animal death and death of cells in culture due to toxicity are similar in that both involve some type of injury. For the animals, the cellular injury leads to tissue and organ injury to the most sensitive target organ, which may then cause the death of the whole organism. An organ system failure can be due either to the death of cells in the affected organ or to the loss of function of the surviving cells in the organ, which results in cell death or loss of function in other organs (GENNARI et al., 2004). The death of an animal is a result of major organ system failure. Ultimately, the cardiovascular and respiratory systems fail. Respiratory depression may be due to the depression of the central nervous system rather than a direct assault on the respiratory system. Other major organ system failures, such as liver and kidney failure, gastrointestinal corrosion, and bone marrow depression, also cause death. Cell death is a culture system that involves the death of a single cell type. Cell death and animal death may be similar since animals are comprised of organ systems consisting of tissues, which are comprised of cells. All cells, regardless of whether they are animal cells or in vitro cell cultures, have similar cellular mechanisms of energy production, utilisation, and maintenance of cell membrane integrity. Animal death and death of cells in culture due to toxicity are similar in that both involve some type of injury. For the animals, the cellular injury leads to tissue and organ injury to the most sensitive target organ, which may then cause the death of the whole organism.

Considering the results obtained in vivo, it is possible that the in vitro NRU cytotoxicity test predicted the toxicity of the A. crassiflora seed extract by estimating a lower LD_{50} value than that necessary to produce lethal death. The endpoint measured in the in vitro NRU cytotoxicity tests is cell death, and the endpoint measured in the acute systemic toxicity assays is usually animal death. Cell death and animal death may be similar since animals are comprised of organ systems consisting of tissues, which are comprised of cells. All cells, regardless of whether they are animal cells or in vitro cell cultures, have similar cellular mechanisms of energy production, utilisation, and maintenance of cell membrane integrity. Animal death and death of cells in culture due to toxicity are similar in that both involve some type of injury. For the animals, the cellular injury leads to tissue and organ injury to the most sensitive target organ, which may then cause the death of the whole organism.

Figure 1. Cytotoxicity evaluation for the cerrado fruits extracts: a) cagaita seed; b) araticum seed; c) araticum peel; and d) pequi peel. The graphs include the original concentration-response data and the curves generated from the bootstrap analysis.

the 14th day. However, rats that received the A. crassiflora seed extract showed some anxiety and were restless, perhaps because of the phenolic acids and their derivatives (caffeic acid, caffeoyl glucose and caffeoyl tartaric) present in this extract. In addition, it was observed that there was significant reduction in the body weight gain (approximately 28 and 23%, respectively) of the two groups that received seed extract if compared with the control group (Roesler et al., 2007a). The same study was conducted for the A. crassiflora peel extract, in which two groups of seven rats each received an aqueous solution of 50 mg of gallic acid equivalents per kg orally, i.e. 358 mg of crude extract per kg of weight for 14 days. No different behaviour was observed for these two groups. Thus, the in vitro result is in agreement with the previous in vivo study once there was no sign of cell viability decrease up to the maximum concentration tested. In addition, the in vitro method was able to estimate that the A. crassiflora seed extract could be harmful since it showed moderate citotoxicity.

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toxin is delivered to cell, and how it is distributed, metabolised, and excreted. The 3T3 cell culture system includes serum and has little or no capacity to metabolise xenobiotic compounds. The excretion from the cell culture milieu cannot occur since cell culture systems have no excretory system. The culture cells are exposed to substances during the test system.

3.2 Phototoxicity

Differently from the in vitro cytotoxicity results that are used to screen toxicity and to estimate the LD₅₀ of chemicals, the NRU phototoxicity assay using 3T3 cells was validated by ECVAM and accepted for regulatory use to detect the phototoxic potential of substances. The phototoxic potential is assessed by comparing the differences in toxicity between negative control plates that were not exposed to UVA and test plates exposed to UVA (Figure 2).

Considering the analysis of the PIF values (Table 3), none of the extracts showed phototoxic potential in the dosage levels tested; only the pequi peel and araticum seed ethanolic extract presented cell viability reduction during the exposure to UVA light. The mean PIF obtained for the pequi peel and seed extracts were 1.0 and 1.6, respectively. Although there was cell viability reduction, it occurred even for the no exposed plates so the PIF value was relatively low and the extracts were not considered phototoxic (PIF < 5). Due to the low solubility of cagaita seed and araticum peel extracts, it was not possible to increase the concentration (it was tested up to 300 µg.mL⁻¹) in order to obtain the EC₅₀ values with or without UVA radiation. Consequently, it was not possible to define their respective PIF values.

The MPE values (Table 3) were also calculated and the non-phototoxic potential was confirmed for the extracts of araticum.

Figure 2. Phototoxicity evaluation for the cerrado fruits extracts: a) cagaita seed; b) araticum seed; c) araticum peel; and d) pequi peel. Gray and white points represent data from dark and light curves, respectively. The graphs include the original concentration-response data and the curves generated from the bootstrap analysis.
seed, cagaita seed, and pequi peel. However, for the araticum peel extract it was found a MPE value of 0.135 indicating a phototoxic potential that could not be predicted using PIF because of the low solubility of this extract.

Table 2 presents some chemicals PIF and MPE values for comparative purposes, and the correspondent p-values calculated in the PHOTOTOX software. The phototoxicity results obtained by the in vitro NRU method are of great importance especially for cosmetic and pharmaceutical application since topical formulations are commonly used during the day with exposure to the sun and artificial light.

4 Conclusions

Brazil has a strong tradition of herbal medicine and, like most developing countries, its rural and traditional population still depends mainly on the indigenous system of medicine for their healthy related matters. Herbal medicines are frequently used to treat a large variety of ailments and symptoms, such as fever, inflammation, and pain. However, there is little information about their efficacy and lack of acute toxicity.

The Annonaceae family has a great variety of exotic fruits, which are apparently rustic and have typical form such as sugar-apple or sweetsop (Annona squamosa), graviola (Annona muricata), and araticum of cerrado or marolo (Annona crassiflora) (SILVA; TASSARA, 2001). Many members of Annonaceae are used in folk medicine for antiparasitic or antitumor treatment of intestinal diseases. In Brazil, the fruits are sold in regional markets and have no commercial value. The seeds in oil are used against scalp infections and in folk medicine; the leaves and seeds infusion is used against diarrhoea and as homemade preparations such as juice, ice cream, and jelly (SILVA et al., 1994). The volatile constituents of the Eugenia dysenterica leaf oil has been investigated for its antifungal activity (COSTA et al., 2000).

In conclusion, the in vitro NRU methods showed that the excellent antioxidant extracts of the araticum seed, cagaita seed, and pequi peel did not present phototoxic hazard; only the araticum peel extract presented a phototoxic potential indicating that complementary studies should be carried out to evaluate the possible in vivo effects if any topical use of this extract is intended. With regard to cytotoxicity, the araticum peel and cagaita seed extracts did not show any inhibition of cell grown up to 300 µg.mL⁻¹ the maximum solubility of these extracts. The araticum seed extract presented an IC₅₀ of 108.0 ± 7 µg.mL⁻¹ and the pequi peel extract had an estimated IC₅₀ of 3731.3 ± 579 µg.mL⁻¹. The LD₅₀ obtained by using the RC rat-only weight regression for the araticum seed extract was 831.6 mg.kg⁻¹ and for the pequi peel extract it was 2840.7 mg.kg⁻¹. While the in vitro NRU phototoxicity method is validated and accepted for regulatory uses, the cytotoxicity data should be used as a screening procedure to assess cytotoxicity as well as one of the factors used to estimated the starting dose for in vivo acute lethality studies based on the preliminary information that this approach could reduce the number of animals used in in vivo studies (i.e. reduction), minimize the number of animals that receive lethal doses (i.e. refinement), and avoid underestimating hazard. A detailed evaluation of tropical plants used in local

### Table 3. Values of phototoxic potential (PIF and MPE) by in vitro NRU and p values.

| Extract or chemical compound | In vivo | PIF | p-value × 100% | MPE | p-value × 100% |
|----------------------------|--------|-----|----------------|-----|---------------|
| Cagaita seed               | ND     | 1.0*| 0%             | 0.09| 21%           |
| Araticum seed              | ND     | 1.6 | 0%             | 0.03| 0%            |
| Araticum peel              | ND     | 1.0*| 0%             | 0.14| 95%           |
| Pequi peel                 | ND     | 1.0 | 0%             | 0.00| 0%            |
| Sodium lauryl sulphate     | Not phototoxic | 1.3 | 1% | 0.01 | 0% |
| Penicillin G               | Not phototoxic | 3.2 | 21% | 0.09 | 29% |
| Terephthalidene dicamphor s.a. | Not phototoxic | 1.6 | 63% | 0.01 | 9% |
| Bergamot oil               | Phototoxic | 15.2 | 0.79 | 0.37 | 0.91 |
| Chlorpromazine             | Phototoxic | 36.0 | 100% | 0.34 | 100% |
| Furosemide                 | Phototoxic | 91.8 | 43% | 0.15 | 54% |

The different compounds in the table are references for phototoxic potential comparison. These data were based on Peters and Holzhütter (2002). ND: not determined. *PIF was considered 1.0 because no EC₅₀ value exists for both curves.
health traditions and the ethnopharmacological evaluation to verify their efficacy and safety can lead to the development of invaluable herbal drugs or isolation of compounds of therapeutical value. It is expected that the search for an economically viable and environmentally correct application of cerrado natural resources, such as natural antioxidants, could increase the value of these fruits and help avoid the destruction of natural vegetation and cerrado biodiversity.

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