Nuclear and mitochondrial genome instability induced by senna (*Cassia angustifolia* Vahl.) aqueous extract in *Saccharomyces cerevisiae* strains

C.R. Silva¹,²,³, A. Caldeira-de-Araújo¹, A.C. Leitão² and M. Pádula¹,³

¹Laboratório de Radio e Fotobiologia, Departamento de Biofísica e Biometria, IBRAG, Universidade do Estado do Rio de Janeiro, Rio de Janeiro, RJ, Brasil
²Laboratório de Radiobiologia Molecular, Universidade Federal do Rio de Janeiro, Rio de Janeiro, RJ, Brasil
³Laboratório de Microbiologia e Avaliação Genotóxica, Faculdade de Farmácia, Centro de Ciências da Saúde (A2-42), Universidade Federal do Rio de Janeiro UFRJ, Rio de Janeiro, RJ, Brasil

Corresponding author: M. de Pádula
E-mail: marcelo@pharma.ufrj.br

Received February 17, 2014
Accepted May 29, 2014
Published November 27, 2014
DOI http://dx.doi.org/10.4238/2014.November.27.13

**ABSTRACT.** *Cassia angustifolia* Vahl. (senna) is commonly used in self-medication and is frequently used to treat intestine constipation. A previous study involving bacteria and plasmid DNA suggested the possible toxicity of the aqueous extract of senna (SAE). The aim of this study was to extend the knowledge concerning SAE genotoxicity mechanisms because of its widespread use and its risks to human health. We investigated the impact of SAE on nuclear DNA and on the stability of mitochondrial DNA in *Saccharomyces cerevisiae* (wt, *ogg1*, *msh6*, and *ogg1msh6*) strains, monitoring the formation of petite mutants. Our results demonstrated that SAE specifically increased Can⁸ mutagenesis only in the *msh6* mutant, supporting the view that SAE can induce misincorporation errors in DNA.
We observed a significant increase in the frequency of petite colonies in all studied strains. Our data indicate that SAE has genotoxic activity towards both mitochondrial and nuclear DNA.

Key words: DNA repair; Genome instability; Oxidative damage; petite colonies; Saccharomyces cerevisiae

INTRODUCTION

Many herbal laxatives, such as Cassia angustifolia Vahl. (senna), have been used for self-medication in the treatment of intestinal constipation (Brusick and Mengs, 1997; Van Gorkon et al., 1999).

In our previous study, we assessed the possible toxicity of senna. We reported that the aqueous extract of senna (SAE) exhibited highly genotoxic activity such as DNA strand breaks against plasmid DNA in a cell-free system (Silva et al., 2008). However, SAE showed antioxidant/antimutagenic activity in Escherichia coli cells upon \( \text{H}_2\text{O}_2 \) treatment. This feature of SAE was observed in the E. coli IC203 (\(uvrA\) oxy\(R\)) and IC205 (\(uvrA\) mut\(M\)) strains, which are highly sensitive to oxidative mutagens (Silva et al., 2008).

As part of a continuous effort to understand the toxic effects of senna, we examined SAE genotoxicity mechanisms in Saccharomyces cerevisiae. S. cerevisiae is an interesting eukaryotic model for genetic studies, as a number of yeast proteins have been shown to be functionally interchangeable with human proteins (Jazayeri and Jackson, 2002; Mager and Winderickx, 2005). Additionally, yeast are useful for understanding base excision repair, genetic recombination, and response to stresses involved in cellular and organism aging (Gershon and Gershon, 2000; Prakash and Prakash, 2000; Rowe et al., 2008).

Oxidative damage produced by intracellular reactive oxygen species results in DNA lesions, as base modifications, and in the formation of apurinic/apyrimidinic lesions, which may be toxic and/or mutagenic. Mutagenic 8-oxo-7, 8-dihydroguanine (8-oxoG) lesions are present in elevated levels in aged and cancerous cells (Rowe et al., 2008).

In S. cerevisiae, mismatch repair (MMR) and translesion synthesis cooperate with Ogg1 to prevent the mutagenic effect of 8-oxoG in nuclear DNA (Girard and Boiteux, 1997; Boiteux et al., 2002; de Pádula et al., 2004). Msh2/Msh6 proteins contribute to the prevention of GC to TA transversions in nuclei, as the Msh6 protein can remove adenine from 8-oxoG:A pairs (Boiteux et al., 2002). In addition, Ogg1 has been implicated in the prevention of petite mutants of S. cerevisiae (Singh et al., 2001), indicating a role of Ogg1 in avoiding 8-oxoG in mtDNA. The Msh1 protein has been implicated in mitochondrial stability, indicating that MMR occurs in mitochondria (Larsen et al., 2005).

In this study, we evaluated the genotoxic potential of SAE using a series of S. cerevisiae strains in 2 experimental assays, including determination of canavanine-resistant mutants and determination of mitochondrial mutants (petite colonies) upon SAE treatment.

MATERIAL AND METHODS

Phytopharmaceutical, chemical agents, and reagents

Orient Mix Fitoterápicos do Brasil, Ltd. (Rio de Janeiro, Brazil) provided powdered...
senna leaves in capsules; each capsule contained 35% sennosides, A and B, according to data from bulla. The SAE in 0.9% NaCl sterile solution was extemporaneously prepared for each experiment (Silva et al., 2008). Bacto agar, bacto-peptone, yeast nitrogen base without amino acids, and bacto yeast extract were purchased from Difco Laboratories (Detroit, MI, USA). Sodium chloride, potassium dihydrogen phosphate, sodium phosphate dibasic dodecahydrate, triphenyltetrazolium chloride, sodium bicarbonate sodium hydroxide, and glycose were obtained from Merck (White House Station, NJ, USA); uracil, L-lysine, L-tryptophan, L-leucine, L-canavanine sulfate salt, and L-histidine monohydrochloride monohydrate were from Sigma-Aldrich (St. Louis, MO, USA). Ultra-pure water was obtained using a Milli-Q water system from Millipore Corporation (Billerica, MA, USA).

**Media and growth conditions**

*Schizosaccharomyces pombe* strains FF18733 (MATa, leu2-3-112, trp1-289, his7-2, ura3-52, lys1-1), CD138 (as FF18733 but ogg1Δ::TRP1), BPS1031 (as FF18733 but msh6Δ::KANMX6), and BPS1050 (as FF18733 but msh6Δ::KANMX6, ogg1Δ::TRP1) were used (Heude and Fabre, 1993; Thomas et al., 1997; Melo et al., 2004; de Pádua et al., 2004). Cells were grown at 30°C in YPD medium (1% yeast extract, 1% bacto-peptone, 2% glucose, with 2% agar for plates) or yeast nitrogen base dextrose (YNBD) medium (0.7% YNBD without amino acids, 2% glucose, with 2% agar for plates) supplemented with appropriate amino acids and bases (de Pádua et al., 2004). Supplemented YNBD medium lacking arginine but containing canavanine (60 mg/L) was used for the selective growth of canavanine-resistant (CanR) mutants (de Pádua et al., 2004).

**Spontaneous and induced mutation frequencies**

Yeast strains were grown in 2 mL YPD medium with for 2 days at 30°C. For each strain, 10 independent cultures in YPD (2.0 mL) were inoculated with approximately 2 x 10^5 cells and grown at 30°C for 2 days with SAE or 0.9% NaCl sterile solution. A concentration of 500 mg/mL SAE was chosen for these experiments based on our previous study in which this concentration was generated DNA strand breaks in plasmid DNA (Silva et al., 2008). Cell density was measured by plating dilutions on YPD agar plates and counting the colonies after 2-3 days of growth at 30°C. The quantification of CanR mutants was determined after plating on selective medium (YNBD-canavanine). All experiments were carried out independently 3 times (Melo et al., 2004). Mutation frequencies were determined as CanR per 10^7 cells (Melo et al., 2004).

**Measurement of mitochondrial mutants**

*Petite* mutants form white colonies (*petite* colonies). Cultures were diluted, plated on YPD, and incubated at 30°C for 3-4 days. Spontaneous and induced mitochondrial mutants were scored using a triphenyltetrazolium chloride color assay (Ogur et al., 1957).

**Statistical analysis**

The results were analyzed by analysis of variance when i) the data were normally
distributed as verified by the Kolmogorov and Smirnov method and ii) samples from the populations showed identical standard deviations as verified using the Bartlett method. Analysis of variance was followed by the Student-Newman-Keuls multiple comparison test using the statistical program InStat version 3.01 (GraphPad Software, San Diego, CA, USA). A significance level of 5% was used to evaluate the data.

RESULTS

SAE-induced mutagenesis in *S. cerevisiae* was assessed using FF18733 (wild-type, *wt*), CD138 (*ogg1*), BPS1031 (*msh6*), and BPS1050 (*msh6ogg1*) strains. SAE was able to specifically increase mutagenesis only in *msh6* cells (*P* < 0.05). In this strain, the number of Can^r^ mutants/10^7^ cells (699) was 6-fold higher than of the rate of spontaneous revertants (116) after 96 h of treatment with SAE, while no significant increase was observed in the other strains tested (*P* > 0.05).

Regarding mitochondria mutagenic damage, SAE increased the number of respiratory mutants in all strains tested. The petite colonies fold-increase over spontaneous levels were *wt*: 2.6; *ogg1*: 3.3; *msh6*: 2.5, and *msh6ogg1*: 1.7 after 96 h of treatment with SAE. These results indicate a distinct genotoxicity pattern of SAE in eukaryotic cells, as nuclei and mitochondria endure different types of DNA damage.

DISCUSSION

In the present study, we investigated the impact of SAE on mutagenesis and mitochondrial stability in *S. cerevisiae* strains.

We evaluated whether SAE was genotoxic in *S. cerevisiae* by monitoring the ability of SAE to induce mutations in *wt, ogg1, msh6*, and *ogg1msh6* strains.

Our results showed that SAE specifically increased Can^r^ mutagenesis only in the *msh6* mutant, supporting that SAE induces misincorporation errors in nuclear DNA, as *MSH6* plays a direct role in MMR, eliminating pre-mutagenic base mismatches (Ni et al., 1999). As no increase in mutagenesis was observed in *ogg1* mutants, the induction of 8-oxoG by SAE can be ruled out, as well as the formation of 8-oxoG:A mispairs in nuclear DNA. Notably, SAE did not increase the number of Can^r^ mutants of the *msh6ogg1* strain. In fact, double inactivation of *OGG1* and *MSH6* genes had a synergistic effect on spontaneous mutagenesis (Radak and Boldogh, 2010). This may compromise the detection of other base substitutions, as approximately 80% of base substitution events are spontaneous GC to TA transversions in this strain (Ni et al., 1999; Radak and Boldogh, 2010). These results reinforce the notion that SAE-induced DNA damage involves other pre-mutagenic mismatches, likely leading to base pair substitutions such as GC to AT, GC to TA, and AT to GC, independently of 8-oxoG generation (Boiteux et al., 2002).

In addition, we observed a significant increase in the frequency of petite colonies induced by SAE treatment (*P* < 0.05). In the *wt* strain, SAE induced a 2.6-fold increase in the number of petite colonies, while a 3.3-fold increase was observed for the *ogg1* strain. This indicates that SAE induced distinct DNA damage profiles according to DNA localization. In the nucleus, SAE-induced DNA damage was a substrate for Msh6 (MMR) but not for Ogg1 protein. In contrast, in mitochondria, DNA damage required Ogg1 protein to prevent SAE-induced petite mutant colonies.
Interestingly, the msh6 mutant displayed a lower level of spontaneous petite mutants compared to the wt strain. Although the Msh6 protein has not been described in mitochondria, MMR (Msh1) has been previously reported to be a backup for mitochondrial base excision repair (Dzierzbicki et al., 2004). Msh6 may interfere with and modulate the generation of petite mutants. As Ogg1 is needed to avoid spontaneous and SAE-induced petite mutants, 8-oxoG appears to be generated in mitochondria. However, this appears to occur in the nucleotide pool rather than in the mtDNA. If 8-oxoG is produced in the nucleotide pool, it can be incorporated in DNA opposite to adenine and can constitute a substrate to Msh6 mismatch repair. Additionally, adenine is eliminated from the 8-oxoG:A mismatch and, after one replication round, may lead to a TA to GC transversion. This renders the wt strain more prone to mitochondrial instability compared to an msh6 mutant. Indeed, the results obtained with the double-mutant ogg1msh6 reinforce this hypothesis. Although inactivation of MSH6 in an ogg1 background was not sufficient for reducing the overall level of spontaneous petite mutants, it was able to partially suppress SAE-induced petite mutants compared to the ogg1 strain. This reinforces that SAE can induce DNA damage to S. cerevisiae in a bimodal manner. While SAE induces nuclear DNA damage requiring mismatch repair but not Ogg1 protein, in mitochondria SAE induces oxidative damage and relies on Ogg1 to prevent petite mutants.

CONCLUSION

Our data indicate a differential genotoxic pattern of SAE in the mitochondria and nucleus regarding DNA damage production.

Conflicts of interest

The authors declare no conflicts of interest.

ACKNOWLEDGMENTS

The authors thank MSc. Marcia B.N. de Oliveira (UERJ), MSc Simone Simplicio (UERJ), MSc Paulo Thiago S. Santos (UERJ), BSc Rita de Cássia Albuquerque (UFRJ) and BSc Janine S.C. Rurr (UFRJ) for their technical assistance; Dr. Alicia V. Pinto (FIOCRUZ) for critical reading of this manuscript. Research supported by CNPq (Proj. Universal - Proc. #474280/2007-1) and FAPERJ (APQ1 #111566/2010) Brazilian agencies.

REFERENCES


