Mitochondrial ATPase activity and membrane fluidity changes in rat liver in response to intoxication with Buckthorn (*Karwinskia humboldtiana*)

Margarita Cid-Hernández1, Ana C Ramírez-Anguiano†1, Genaro G Ortiz‡2, Eddic W Morales-Sánchez‡2, Luis J González-Ortiz†1, Sandra F Velasco-Ramírez†1 and Fermín P Pacheco-Moises*1

Abstract

**Background:** *Karwinskia humboldtiana* (Kh) is a poisonous plant of the rhamnacea family. To elucidate some of the subcellular effects of Kh toxicity, membrane fluidity and ATPase activities as hydrolytic and as proton-pumping activity were assessed in rat liver submitochondrial particles. Rats were randomly assigned into control non-treated group and groups that received 1, 1.5 and 2 g/Kg body weight of dry powder of Kh fruit, respectively. Rats were euthanized at day 1 and 7 after treatment.

**Results:** Rats under Kh treatment at all dose levels tested, does not developed any neurologic symptoms. However, we detected alterations in membrane fluidity and ATPase activity. Lower dose of Kh on day 1 after treatment induced higher mitochondrial membrane fluidity than control group. This change was strongly correlated with increased ATPase activity and pH gradient driven by ATP hydrolysis. On the other hand, membrane fluidity was hardly affected on day 7 after treatment with Kh. Surprisingly, the pH gradient driven by ATPase activity was significantly higher than controls despite an diminution of the hydrolytic activity of ATPase.

**Conclusions:** The changes in ATPase activity and pH gradient driven by ATPase activity suggest an adaptive condition whereby the fluidity of the membrane is altered.

**Keywords:** *Karwinskia humboldtiana*, ATPase, Mitochondria, Membrane fluidity, pH gradient

Background

*Karwinskia humboldtiana* currently known by the common names buckthorn, wild cherry, tullidora, capulin tullidor, capulincillo, coyotillo, and cacatsin, is a poisonous shrub of the Rhamnacea family that grows from Southern United States to northern Colombia [1]. Medicinal properties have been attributed to several species of the genus *Karwinskia* due to a secondary metabolite with antineoplastic action on mammalian tumor cells [2].

The ingestion of the green or ripe fruit of Kh causes a flaccid, symmetric, progressive, and ascending palsy of the lower limbs, which, in severe cases may progress to quadriplegia and breathing insufficiency, with victims requiring assisted mechanical ventilation. The neurologic symptoms are similar to those of poliomyelitis, Guillain-Barre syndrome, and other polyradiculoneuritis syndromes, for which it is often mistaken. In Mexico, the accidental ingestion of Kh causes several health problems [3-5], particularly there are numerous reports of poisoning by Kh, of which 68.1% of cases occurred in children under 5 years old [5]. Kh contains significant amounts of phenolic antracenonas and/or anthraquinone [3,6].

Oral administration of a slurry prepared with the seed of Kh in rats causes a uncoordinated motor activity of the hind limbs followed by rapidly ascending flaccid paralysis and signs of respiratory distress [7]. A histological
study showed lesions in peripheral nerves corresponding to a segmental demyelinating neuropathy and specific lesions in the cerebellum, cerebral cortex, reticular formation, possibly due to anoxia [8]. Deaths in cats have been reported after acute administration of higher doses [9]. Paralysis caused by Kh is due to blockage in the conduction of nerve impulses, and to muscle denervation [7]. Other authors have reported damage in organs such as liver (necrosis and fatty degeneration), lung (pulmonary hemorrhage), heart (necrosis) and kidney [10]. Furthermore, a decrease of peroxisomes has been reported in rat hepatocytes [11].

The structure and function of cell membranes plays an important role in maintaining intracellular homeostasis through the activity of membrane enzymes, hormone receptors, and the transmembrane transport system. These dynamic phenomena are influenced by membrane fluidity. Fluidity is an important characteristic of biologic membranes at a molecular level. It indicates the way and the rate of motion of molecules in the membrane, and is inversely related to membrane microviscosity. Changes in membrane fluidity may play a role in the regulation of membrane properties, both under normal and pathological conditions [12]. Mitochondria provide cellular energy via oxidative phosphorylation, using the multisubunit complexes of the respiratory chain to create a transmembrane potential, which drives ATP synthesis by the ATP synthase; ATP synthase is also capable of working as an ATPase [13].

It has been reported that mice fed with Kh showed some hepatic changes such as degeneration and necrosis, in addition to neurologic disease and weight loss [14]. Furthermore, a single dose of an oral preparation from the seed fruits of Kh causes a significant decrease of ATP concentrations in renal tissue [10] and liver [15]. In mitochondria isolated from liver of rats, the petroleum ether extracts from Kh fruits, induces inhibitory and coupling effects on respiration and oxidative phosphorylation [16], suggesting that alterations on energetic metabolism and oxidative stress are related with the mechanism by which Kh induces its cytotoxic effects. These findings prompted us to investigate the effect of an extract of Kh seeds on membrane fluidity and mitochondrial ATPase activities in rat liver.

Methods

Plant material
Fruits were collected from August to October, 2010 at the settlements of Santa Cruz and Santa Gertrudis, municipality of Colotlán, state of Jalisco (México) and were dried at room temperature and light protected. Before its administration, the almond and cuticle of the seeds were removed, then ground, weighed and mixed with peanut butter to form a homogeneous paste (1:3, weight/weight).

T-514 anthracenone from the cuticle and almond of the seeds of K. humboldtiana were isolated and quantified by procedures already described [17]. The content of T-514 in the samples used was 20.5% of the total anthracenones. (1 g of tissues contains 0.13 g of toxin T-514).

Experimental animals
Thirty five adult male Sprague Dawley rats (obtained from the animal facility of Centro de Investigación Biomédica de Occidente, Guadalajara, Jalisco, Mexico) weighing 250–300 g, maintained on a 12 h/12 h light–dark cycle at 25 ± 2°C, fed with standard diet (Cho Purina) and water ad libitum were used for this research work. All animals received humane care based on the international guidelines on the ethical use of animals, and the “Norma Oficial Mexicana NOM-062-ZOO-1999”. Experimental procedures were approved by the local animal ethics committee (Guadalajara, Jalisco, Mexico). Animals were randomly assigned into control non-treated group (n = 5) that received only vehicle (peanut butter) and three Kh treated groups (n = 10) that received 1, 1.5 and 2 g/Kg body weight of dry powder of Kh fruit, respectively. We take special care to ensure that all peanut butter was eaten by rats. Five rats of each group were euthanized at day 1 and 7 after treatment, respectively. Rats euthanized at day 7 received a second dose of Kh, 24 h after the first dose.

Submitochondrial particles isolation
After treatment, rats were sacrificed by decapitation and livers were immediately dissected for mitochondria isolation. Liver was homogenized with a teflon-on-glass homogenizer Potter-Elvejhem in 35 ml cold buffer containing 250 mM sucrose, 10 mM 4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid (pH 7.4), 1 mM ethylene glycol tetraacetic acid, and kept on ice. The homogenate was centrifuged at 600 x g for 5 min at 4°C. The pellet was discarded, and the supernatant was centrifuged at 8,000 x g for 10 min at 4°C. The foamy layer at the top of the supernatant was removed. The mitochondrial pellet was washed with buffer containing 0.1% fatty acid-free serum albumin and finally resuspended in buffer to contain about 20 mg/ml protein. Then, submitochondrial particles were obtained as previously described [18]. Membrane protein was measured by the method of Lowry et al. [19] using bovine serum albumin as standard.

Membrane fluidity estimations
The degree of membrane lipid fluidity was quantitatively estimated from the excimer to monomer fluorescence
intensity ratio (Ie/Im) of the fluorescent probe 1,3 dipyr-
enylpropane (DPP) incorporated in submitochondrial
particles. Briefly, 0.25 mg of membranal protein and
0.1 nmol DPP were mixed with 10 mM Tris–HCl buffer
(pH 7.8). The mixtures were incubated in darkness at 4°C
for 3 hours, in order to achieve maximal incorporation of
the fluorescent probe to the membranes. Fluorescence
was measured at 30°C on a Perkin Elmer fluorescence
spectrometer, LS50B. The fluorophore was excited at
329 nm and the monomer and excimer fluorescence
intensities were read at 379 and 480 nm, respectively.
From these readings, the Ie/Im was calculated. Fluores-
cence corrections obtained from readings of mem-
branes without DPP were applied to all fluorescence
values [20].

ATPase activity quantification
Oligomycin-sensitive ATPase rates were assayed by the
release of inorganic phosphate. The standard reaction
medium (0.5 mL) contained 125 mM KCl, 40 mM Hepes/KOH (pH 8.0), 0.1 mM EGTA, 3 mM ATP,
5 mM MgCl₂ and 2.2 mM oligomycin. The reaction was
initiated by the addition of submitochondrial particles
(0.25 mg of protein) [21]. It was quenched with 200 mL
of cold 30% (w/v) trichloroacetic acid. Afterwards, the
sample was centrifuged for 10 min at 3,500 rpm and
400 μL of the supernatant was separated, and 1 mL of
3.3% ammonium molybdate was added, followed by
100 μL of 10% ferrous sulphate. The reading of the ab-
sorbance of the samples was recorded at 660 nm [22]
using the Benchmark Plus Microplate Spectrophotom-
eter System, 110/230 V by Bio-Rad. These analyses were
referred to phosphate curve for estimating the enzyme
activity in submitochondrial particles. Control experi-
ments showed that 2 μM oligomycin inhibits 90–95 of
the ATPase activity in submitochondrial particles.

pH gradient driven by ATPase
The pH gradient driven by mitochondrial ATPase activ-
ity was determined by the fluorescence quenching of
1 μM 9-amino-6-chloro-2-methoxyacridin (ACMA).
ACMA fluorescence quenching is directly related to pH
gradient, i.e., the higher values of fluorescent quenching,
the higher values of pH gradient [23]. Briefly, submito-
chodrial particles (1 mg of protein) were incubated at
37°C in a medium (2 mL) containing 125 mM KCl,
20 mM MOPS (pH 7.5), 5 mM MgCl₂, 0.1 mM EGTA,
1 μM ACMA, 3 mM ATP and 5 mM inorganic phos-
phate. After stabilization of the signal, membranes were
energized with 3 mM ATP. ATPase activity pump pro-
tons into the interior of membranes, acidifying the
internal space and the fluorescent amine were internal-
ized. Therefore a substantial quenching of ACMA fluo-
rescence is detected, indicating that a pH gradient (acidic
inside) was established by the ATPase. In our experi-
mements, addition of 1 nM of nigericin (H⁺/K⁺ exchanger)
restored the signal, demonstrating that the pH gradient
across the membranes resulted from the transport of
protons.

Statistical analysis
Results are expressed as the standard error of the mean
(mean ± SEM). The significance of differences between
groups was evaluated using One-way analysis of variance
(ANOVA) and Bonferroni’s Multiple Comparison Test.
Differences with P < 0.05 were regarded as significant.

Results
To assess the acute effects of Kh on membrane fluidity
and ATPase activities, from each group, half of the rats
were euthanized on day 1 and the remaining half at day
7 post Kh treatment. Figure 1 depicts the relative mem-
brane fluidity of submitochondrial particles through the
relative excimer/monomer (Ie/Im) ratio of DPP; the
higher the Ie/Im ratio, the more fluid is the membrane.
At day 1 of Kh treatment, the dose of 1 and 1.5 Kg/g in-
creases significantly the fluidity of membranes relative to
the control group. However, higher dose (2 g/Kg) does
not modify substantially the fluidity of mitochondrial
membranes. At day 7, membrane fluidity values in Kh-
treated rats were similar to those of controls and no sta-
tistically significant differences were found.

Several major functions of membranes such as enzyme
activities and ligand-receptor interactions depend on
membrane fluidity. Therefore, to determine whether an
alteration in enzymatic activity was related to changes in membrane fluidity, in this work we determined the ATPase activities in submitochondrial particles. A basal ATPase activity was detected in control rats. As shown in Figure 2, administration of Kh induced a significant increase of mitochondrial ATPase activity at the dose of 1 and 1.5 g/Kg, as compared to control rats, on day 1 after Kh treatment. Conversely, ATPase activity showed a significant reduction at day 1 when using a dose of 2 Kg/g, and day 7, at all doses.

Transmembrane pH gradient driven by ATPase activity has been extensively used as indication marker of the correct function of the enzyme proton channel and coupling between transport and catalysis [21,24]. Control experiments showed that ACMA fluorescence quenching was not detected when membranes were incubated briefly with 2 μM oligomycin (Figure 3). A substantial pH gradient driven by ATPase activity was detected in liver submitochondrial particles from non-treated rats. In addition, Kh treatment induces a significant increase of pH gradient at lower doses on day 1 and conversely, at the higher dose tested induces a diminution of pH gradient. To our surprise, pH gradient was higher than control group, on day 7 post-treatment, at all doses tested (Figure 4).

Discussion and conclusions
In this work we found that rats under Kh treatment at all dose tested does not developed any neurologic symptoms (not shown). However, we detected alterations in membrane fluidity and ATPase activity in liver submitochondrial particles. These changes were not linear over the dose-range studied. For instance, lower dose of Kh...
on day 1 after treatment induced higher membrane fluidity than control group. This change was strongly correlated with increases in the apparent ATPase activity and pH gradient driven by ATP hydrolysis. On the other hand, at the higher dose used of Kh extract (2 g/Kg) and day 1 post-treatment, there is no change of the membrane fluidity, but there is a decrease of ATPase activity and therefore a diminished pH gradient. Mitochondrial membrane fluidity was hardly affected on day 7 after treatment with Kh. In addition, the pH gradient driven by ATPase activity was significantly higher than controls despite a diminution of the hydrolytic activity of ATPase. These results suggest an adaptive condition whereby the fluidity of the membrane is altered. It has been proposed that changes in membrane fluidity can affect the conformation and activity of membrane-bound enzymes. However, in this work we found that the relationship between membrane fluidity and ATPase activity remains elusive, particularly at day 7 after Kh treatment.

It has been reported that the anthracenone T-514 isolated from *Karwinskia humboldtiana* showed a homogeneous distribution on subcellular fractions prepared from rat liver treated with an acute dose of T-514, indicating that T-514 can pass easily through subcellular compartment membranes. In addition Kh treatment induced a significant increase of protein in liver homogenates and NADPH-cytochrome P450 reductase mesosomal activity [25]. In consonance with these reported findings we showed that Kh treatment has specific effects on liver mitochondria and to our surprise Kh increases the pH gradient driven by ATPase activity.

The changes of mitochondrial membrane fluidity properties may have clinical relevance and can precede other alterations reported previously, such as the significant decrease of the number of peroxisomes present in the hepatocytes [26], the increased plasmatic activities of aspartate aminotransferase and alanine aminotransferase and decreased liver ATP concentration [15].

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
FPPM, MCH, and GGO carried out conception and design of the study, acquisition of data, analysis and interpretation of data, drafting the manuscript and revising. ACRA, EWMS, LKGO, and SFVR carried out acquisition of data and analysis of data. All authors read and approved the final manuscript.

Author details
1Departamento de Química, Centro Universitario de Ciencias Exactas e Ingenierías, Universidad de Guadalajara, Blvd. Marcelino García Barragán 1421, 44430 Guadalajara, Jalisco, Mexico. 2División de Neurociencias, Laboratorio de Desarrollo, Envejecimiento y Enfermedades Neurodegenerativas, Centro de Investigación Biomédica de Occidente (CIBIO), Instituto Mexicano del Seguro Social (IMSS), Sierra Mojada 800, 44340 Guadalajara, Jalisco, Mexico.

Received: 20 May 2014 Accepted: 27 February 2015 Published online: 19 March 2015

References
1. Knight AP, Walter RG. Other plant affecting the nervous system. In: Knight AP, Walter RG, editors. Guide to Plant Poisoning of Animals in North America. Jackson, Wyoming: Teton New Media; 2001. p. 218–49.
2. Waksman DE, Torres N, Santoyo A, Ramírez R, Piñeiro-López A. Cuantificación de T-514 (Peroxisomicina A1) en 2 plantas del género Karwinskia. Polibotánica. 1997:5–9.
3. Becerra-Verdin EM, Bermúdez-Barba MV, Salazar-Leal ME, Ancor Rodríguez J, Romero-Díaz V, Soto-Domínguez A, et al. Karwinskia humboldtiana (buckthorn) fruit causes central nervous system damage during chronic intoxication in the rat. Toxicon Off J Int Soc Toximology. 2009;53:645–51.
4. Charlton KM, Pierce KRA. Neuropathy in Goats Caused by Experimental Coyotillo (Karwinskia humboldtiana) Poisoning: II. Lesions in the Peripheral Nervous System – Teased Fiber and Acid Phosphatase Studies. Pathol Vet. 1970;7:385.
5. Nava Areolea EM, Vázquez Castañeda JL, González Castañeda ME. Factores geográficos en la epidemiología de la intoxicación por Karwinskia (tullidora) en México. Cad Saude Publica. 2000;16:255–60.
6. Garaz-Ocaras L, Jiang T, Acosta D, Torres-Allans Q, Waksman De Torres N, Piñeiro-López A. Comparison of the hepatotoxicity of toxin T-514 of Karwinskia humboldtiana and its diastereoisomer in primary liver cell cultures. Toxicon Off J Int Soc Toximology. 1994;32:1287–91.
7. Salazar-Leal ME, Flores MS, Sepulveda-Saavedra J, Romero-Diaz VJ, Becerra-Verdin EM, Tamez-Rodriguez VA, et al. An experimental model of peripheral neuropathy induced in rats by Karwinskia humboldtiana (Buckthorn) fruit. Peripheral Nerv Syst. 2006;1:233–61.
8. Ocampo-Rosos LV, Ontiveros-Nieves PG, Fernández-Lucio O. Intoxication with buckthorn (Karwinskia humboldtiana): report of three siblings. Pediatr Dev Pathol. 2007;10:66–8.
9. Waksman N, Martínez L, Fernandez R. Chemical and toxicological screening in genues Karwinskia (Mexico). Rev Latinoam Química. 1989;20:27–32.
10. Jaramillo-Juárez F, Rodríguez VML, Castillo CMQ, Quezada TT, de la Cerda GE, Del Posadas RFA, et al. Daño hepático y en la coagulación de la sangre producido por la administración de los frutos maduros de la planta tallidora (Karwinskia humboldtiana) en la rata. Rev Mex Ciencias Farm. 2009;40:28–34.
11. Sepulveda-Saavedra J, Bermúdez De Rocha MV, Tamez-Rodriguez VA, Ballesteros-Elizondo RG, Moreno-Sepúlveda M, Piñeiro-López A. Quantitative analysis of liver peroxisomes in rats intoxicated with peroxisomicine-A1. Toxicol Lett. 1998;98:71–5.
12. Schuller A, Solís-Henruz JA, Moscat J, Fernández-Queca JC, Municio AM. The fluidity of liver plasma membranes from patients with different types of liver injury. Hapatology. 1986;6:714–7.
13. Boyer PD, The ATP. Synthase-a splendid molecular machine. Annu Rev Biochem. 1997;65:717–49.
14. Bermúdez MV, González-Sánchez D, Guerrero M, Waksman N, Piñeiro A. Experimental intoxication with fruit and purified toxins of buckthorn (Karwinskia humboldtiana). Toxicon Off J Int Soc Toximology. 1986;24:1001–7.
15. Jaramillo-Juárez F, Rodríguez-Vázquez ML, Muñoz-Martínez J, Quezada-Tristán T, Posadas Del Rio FA, Llamas-Viamontes J, et al. The ATP levels in kidneys and blood are mainly decreased by acute ingestion of tallidora (Karwinskia humboldtiana). Toxicon Off J Int Soc Toximology. 2005;46:99–103.
16. Wheeler MH, Camp BJ. Inhibitory and uncoupling actions of extracts from Karwinskia humboldtiana on respiration and oxidative phosphorylation. Life Sci 2 Biochem Gen. Mol Biol. 1971;4:1041–51.
17. Dreyer DL, Arai I, Bachman CD, Anderson WR, Smith RG, Davies GD. Toxins causing noninflammatory paralytic neuropathy. Isolation and structure elucidation. J Am Chem Soc. 1975;97:4985–90.
18. Thayer WS, Rubin E. Effects of chronic ethanol intoxication on oxidative phosphorylation in rat liver sub mitochondrial particles. J Biol Chem. 1979;254:7717–23.
19. Lowy OH, Rosebrough NI, Farr AL, Randall RJ. Protein measurement with the Folin phenol reagent. J Biol Chem. 1951;193:265–75.
20. Ortiz GG, Pacheco-Moises F, El Hafidi M, Jiménez-Delgado A, Macías-Islas MA, Rosales Corral SA, et al. Detection of membrane fluidity in sub mitochondrial particles of platelets and erythrocyte membranes from Mexican patients with Alzheimer disease by intramolecular eximer formation of 1,3 dipropyrophen. Dis Markers. 2008;24:151–6.
21. Martinez Cano E, Ortiz GG, Pacheco-Moises F, Macías Islas MA, Sanchez-Nieto S, Rosales Corral SA. Functional disorder of F0F1-ATPase in submitochondrial particles obtained from platelets of patients with probable Alzheimer disease. Rev Neurol. 2005;40:611–8.
22. Sumner JB. A method for the colorimetric determination of phosphorus. Science. 1964;100:413–4.
23. Rottenberg H, Moreno-Sanchez R. The proton pumping activity of H+ --ATPases: An improved fluorescence assay. Biochim Biophys Acta. 1993;1183:161–70.
24. Baracca A, Barogi S, Carelli V, Lenaz G, Solaini G. Catalytic activities of mitochondrial ATP synthase in patients with mitochondrial DNA T8993G mutation in the ATPase 6 gene encoding subunit a. J Biol Chem. 2000;275:4177–82.
25. Guerrero-Olazarán M, Vlader-Salvadó JM. Natural anthracenone subcellular distribution and effects on NADPH-cytochrome P450 reductase microsomal activity. Drug Chem Toxicol. 1996;19:301–12.
26. Vargas-Zapata R, Torres-González V, Sepúlveda-Saavedra J, Piñeyro-López A, Rechinger KB, Keizer-Gunnink I, et al. Peroxisomicine A1 (plant toxin-514) affects normal peroxisome assembly in the yeast Hansenula polymorpha. Toxicon. 1999;37:385–98.