

Proteomic analysis of non-toxic *Jatropha curcas* byproduct cake: Fractionation and identification of the major components

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ABSTRACT

Jatropha curcas non-toxic genotypes have been reported in Mexico and the press-cake, after oil extraction, represents a potential of new source of protein for food and feed uses. However, the characterization of the press-cake proteins is still unknown. The aim of this work was to carry out the molecular characterization of *J. curcas* seed storage proteins. Proteins in press-cake were pre-fractionated according to the classical Osborne procedure. Main protein fraction in *J. curcas* cake was represented by glutelins, the electrophoretic analysis showed that glutelins and globulins have the same profile, indicating that oil extraction process could have effect on globulins agglomeration. Protein fractions were analyzed by two-dimensional gel electrophoresis and mass spectrometry, results provide a new dataset of protein species or proteoforms that are accumulated in *J. curcas* endosperm. The identification of toxic proteins such as curcin in the non-toxic variety could represent that this protein have important roles in seeds. Regulatory proteins such as proteasome subunits and 14-3-3 were identified. A group of different heat shock and stress defense protein species was detected. Proteases related with inhibitory activity against DPPIV were also detected; this could support the potential use of *J. curcas* cake as nutraceutical food.

1. Introduction

Jatropha curcas L. is a stem-succulent tree native to America, which as *Ricinus* and *Manihot* belongs to the family *Euphorbiaceae*. *J. curcas* was spread by Portuguese sailors to Africa and Asia (Fairless, 2007; Maghuly and Laimer, 2013). This subtropical plant can grow through different climatic and soil conditions, it has several uses in different agricultural systems such as barriers against wind and soil erosion as well as a source of firewood (Dias et al., 2012; Maghuly and Laimer, 2013). *J. curcas* main attraction is due to its high potential as plant for biodiesel production (Maghuly and Laimer, 2013). *J. curcas* seeds contains 30–45% toxic oil, with a high amounts of oleic and linoleic acids which make it more suitable for fuel purposes, as compared with other vegetable oils, due to its high rate fuel consumption and its higher oxidation stability (Fairless, 2007; Gübitz et al., 1999; Openshaw, 2000; Pramanik, 2003). *J. curcas* oil has therefore been used for long time as a raw material for paints and soap production as well as for lamp and lubricating oils (Kumar and Sharma, 2008). Besides, different parts of *J.*

curcas plant contain a range of interesting metabolites and bioactive compounds, which is taking great attention for research as medicinal plant (Sabandar et al., 2013).

The mature seeds of *J. curcas* have a tick endosperm representing more than 90% of their weight in which is embedded a small embryo (Liu et al., 2009). The seeds are rich on protein (25–30%) and oil (55–62%), values that depends on the region agroclimatic characteristics. After oil extraction from dehulled kernels, the oil is converted into biodiesel and the resulting press-cake, which is rich of protein (60–63%), could be an excellent protein source. The proteins present in the press-cake are rich of essential amino acids even with higher values (except lysine) than those reported by the Food and Agriculture Organization reference protein (Haas and Mittelbach, 2000; Martínez-Herrera et al., 2012). However, due to the highly toxic and anti-nutritional compounds presents in *J. curcas* seeds, make the press-cake and oil unsuitable for the use as feedstuff or for human consumption (Maghuly and Laimer, 2013).

In Mexico, *J. curcas* L. grows wild in semitropical and tropical

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climates, however only Mexico has reported a nontoxic *J. curcas* genotype (He et al., 2011; Perea-Domínguez et al., 2017). The seeds of this nontoxic genotype are traditionally used to prepare a range of traditional dishes in Veracruz, Puebla, and Hidalgo States of Mexico (Makkar and Becker, 2009; Martínez-Herrera et al., 2012).

The exploitation of *J. curcas* press-cakes has been limited by the little knowledge about some aspects of the biochemistry of these seeds such as the type and functionality of proteins concentrated in the press-cake. Therefore, it is necessary to generate information on these aspects that would help to design innovative biotechnological approaches for the use of the high amount of proteins present in press-cake for future applications in the food industry.

The availability of the *J. curcas* genome and transcriptome data reported in the public databases (Costa et al., 2010; King et al., 2011) have allowed the increase of studies toward the seed proteome characterization. Shah et al. (2015) have used the label-free quantitative proteome analysis in order to analyse the proteins presents in the whole *J. curcas* seed endosperm in searching for information on phorbol esters biosynthesis, mechanism which still is not elucidated. Pinheiro et al. (2013) and Soares et al. (2014) focused in the proteomic analysis of the inner integument from developing seeds, while Shah et al. (2016) carried out the proteome analysis on gerontoplasts isolated from the inner integument of developing seeds. Liu et al. (2015) carried out the proteomics analysis of oil body associated protein species by using gels-based proteomic technique, whereas Liu et al. (2013) used the comparative proteomic approach to profile the protein changes during seed development.

However, to date there is no information about the physical properties of *J. curcas* seed storage proteins presents in press-cakes. It is well known that the functional and molecular characterization of protein-enriched fractions is the first step in designing strategies that allow them to be integrated as additives or food ingredients (Rezig et al., 2013). In this sense, Osborne's protein classification (Osborne, 1908), which is based on protein solubility characteristics, is the most common method for seed storage protein characterization and seems to be an excellent pre-fractionation step towards proteome characterization. For these reasons the aim of the present work was to carry out the characterization of proteins present in press-cake obtained after oil extraction of non-toxic *J. curcas*. Gel-based proteomics and LC-MS/MS were used in order to obtain a new description of *J. curcas* press-cake proteins; such valuable information will help in the way to design technologies for development of new food products.

2. Material and methods

2.1. *J. curcas* press-cake preparation

Non-toxic *Jatropha curcas* seeds (Puebla's ecotype, voucher specimen numbered 53203) were obtained from ripe fruits from plants cultivated in "La Esmeralda" Experimental Field of the Agriculture Association of West Sinaloa River (AARSP, Guasave, Mexico). *J. curcas* seed press cake was produced after oil pressing from dehulled seeds (kernels) using a screw-press. Two press cake samples (500 g each) were collected and air-dried under ambient conditions. The combined air-dried press cakes were milled to a small particle size (40-mesh) using a Cyclotec 1093 mill (FOSS Tecator, Hilleroed, Denmark) and then defatted with *n*-hexane (1:10 w/v) for 4 h with agitation and air-dried in a fume hood after decantation of the hexane. Dry meals were kept in plastic bags at 4 °C until used.

2.2. Meal protein sequential fractionation based on Osborne's solubility

Protein fractions were sequentially extracted according to a modified Osborne's procedure (Ribeiro et al., 2004). Briefly, the defatted meal was extracted with water containing 10 mM CaCl₂ and 10 mM MgCl₂ (1:10 w/v) pH 8.0, with constant stirring for 4 h at 4 °C.

The slurry was centrifuged at 30,000g for 1 h at 4 °C. The supernatant was recovered and extensively dialyzed (12.4 kDa MW cut-off) against distilled water. After dialysis, the suspension was centrifuged at 15,000g for 15 min at 4 °C and supernatant (albumins) was collected and freeze-dried.

The resulting pellet from albumins extraction was resuspended with 100 mM Tris-HCl, pH 7.5, containing 10% NaCl (w/v), 10 mM ethylenediaminetetraacetic acid (EDTA), and 10 mM ethylene glycol-bis(2-aminoethylether)-N,N,N',N'-tetraacetic acid (EGTA) (1:10 w/v). The suspension was stirred for 4 h at 4 °C. The insoluble proteins were removed by centrifugation at 30,000g for 1 h at 4 °C. The supernatant (globulins fraction) was dialyzed, centrifuged and freeze-dried.

The insoluble pellet resulting from globulins extraction was resuspended in 75% aqueous ethanol (1:10 w/v) and centrifuged (30,000g at 4 °C for 15 min). The supernatant was dialyzed against distilled water. The supernatant (prolamins) was recovered and freeze-dried. The insoluble pellet from the previous extraction was resuspended (1:10 w/v) with 50 mM sodium borate buffer, pH 10, containing 1% (v/v) β-mercaptoethanol, and 1% (w/v) sodium dodecyl sulphate (SDS). The suspension was stirred at room temperature for 2 h and centrifuged at 30,000g for 15 min at 20 °C. After dialysis against distilled water, the fraction (glutelins) was freeze-dried. All samples were kept at -70 °C until use. A scheme of the whole extraction procedure is presented in Supplementary Fig. S1.

2.3. Protein sample preparation

Freeze-dried protein fractions (five grams) were mixed with 50 mL of its respective extraction solution and 1% (w/v) of polyvinylpyrrolidone (PVPP). The suspensions were mixed in a vortex for 1 min and sonicated for 2.5 min at 20 kHz with 35% of amplitude (GE-505, Ultrasonic Processor, Sonics & Materials, Inc., Newtown, CT, USA). Samples were maintained at 4 °C during suspension steps and centrifuged at 15,000g for 15 min at 4 °C (Beckman Avanti J26-XP, Beckman Coulter, Brea, CA, USA). Supernatants were transferred to new tubes and proteins were precipitated by adding four volumes of ice-cold acetone and incubated overnight at -20 °C. After 15 min of centrifugation at 15,000g for 15 min at 4 °C, the supernatant was decanted and discarded, with the residual pellet being washed twice with ice-cold acetone and allowed to dry at room temperature. The resulting dried pellets were suspended in rehydration buffer (8 M urea, 2% (w/v) 3-[(3-cholamidopropyl)-dimethylammonio]-1-propanesulfonate (CHAPS), 0.56% (w/v) dithiothreitol (DTT), 0.002% bromophenol blue) mixed in vortex for 30 s and sonicated for 80 s. The suspensions were centrifuged under the previous conditions and the supernatants were recovered. Protein concentration was determined by using protein assay (Bio-Rad, Hercules, CA, USA) with bovine serum albumin (BSA) used as standard.

2.4. Two-dimensional gel electrophoresis (2-DE) and gel image analysis

Proteins (900 µg) were suspended in 450 µL of rehydration buffer, containing 0.5% (v/v) IPG buffer (Bio-Rad) in the range of 5–8 (albumins) and 3–10 for globulins and glutelins fractions. Proteins were loaded onto 24 cm linear immobilized pH gradient (IPG, Bio-Rad) strips 5–8 (albumins) or 3–10 (globulins and glutelins). Passive rehydration was carried out at room temperature for 14–16 h. The IEF was conducted at 50 mA per IPG strip and 20 °C in an Ettan IPGphor 3 system (GE Healthcare, Piscataway, NJ, USA) under the following conditions: (I) 150 V gradient for 2 h (II) 200 V gradient for 2 h, (III) 400 V gradient for 2 h, (IV) 1500 V gradient for 2 h, (V) 4500 V gradient for 3 h, (VI) 10,000 V gradient for 3 h; and (VII) holding at 10,000 V for 10 h. After IEF, the IPG strips were stored at -20 °C or immediately equilibrated for 15 min in equilibration buffer [50 mM Tris-HCl pH 8.8, 6 M Urea, 30% (v/v) glycerol, 2% (w/v) SDS, 0.002% (w/v) bromophenol blue, 65 mM DTT].

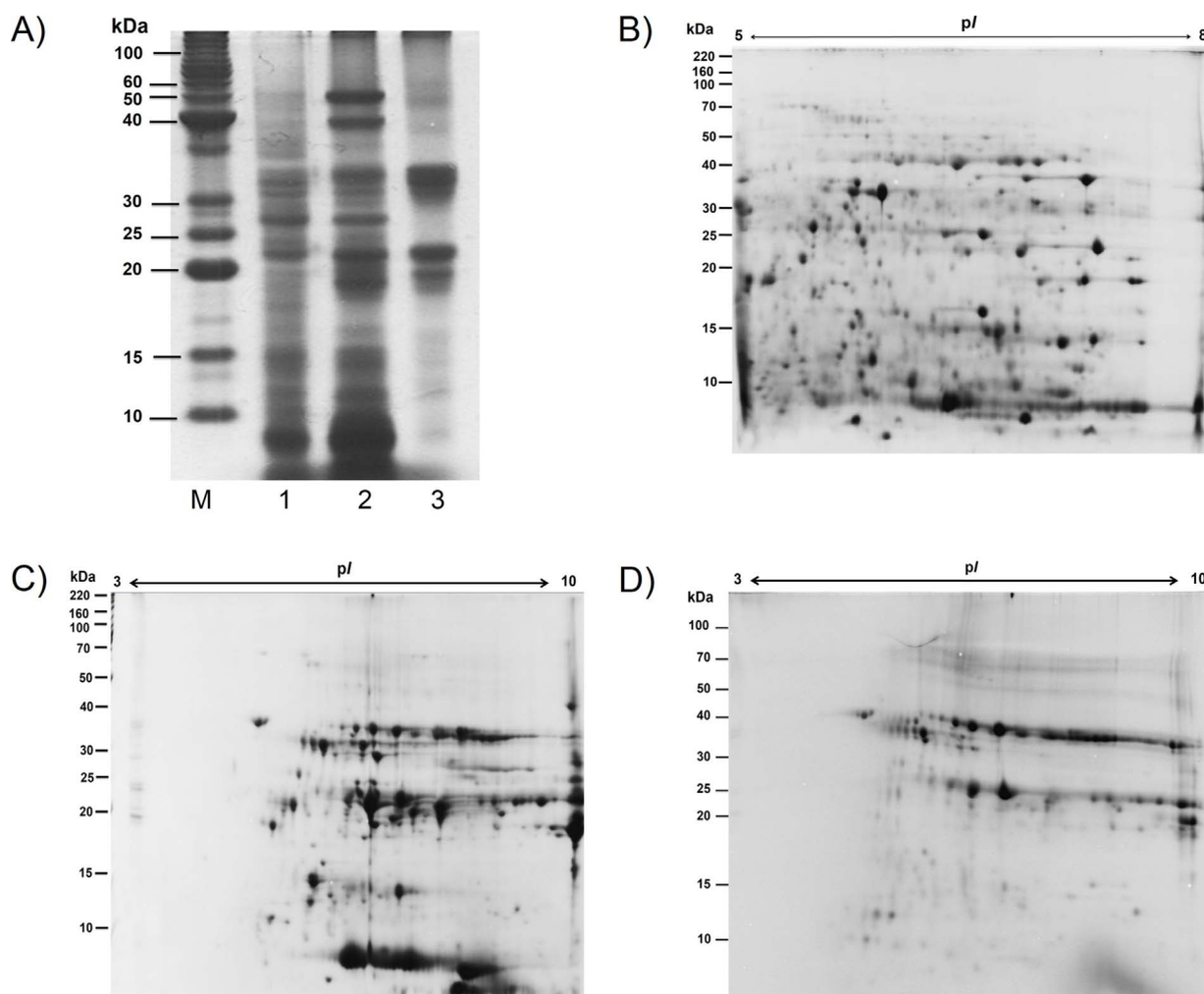


Fig. 1. A) One-dimension electrophoretic profile of *Jatropha curcas* seed storage proteins. Line M = molecular weight markers, Line 1 = albumins, Line 2 = globulins, Line 3 = glutelins. Representative 2-DE gels of *J. curcas* seed storage proteins for B) albumins; C) globulins; D) glutelins.

Strips were placed directly onto 13% polyacrylamide-SDS slab gels and the second dimension was performed using an Ettan™ DALT-six Electrophoresis Unit (GE Healthcare), using SDS electrophoresis buffer [25 mM Tris pH 8.8, 192 mM glycine, and 0.1% (w/v) SDS] and resolved at 20 mA/gel until the dye (bromophenol blue) reached the bottom of the gels. Three different extractions were prepared for gel replicates using the same method. After SDS-PAGE gels were stained with PhastGel™ Blue R-350 (GE Healthcare) and scanned at 100 μ m resolution using a Pharos FX Plus Molecular Imager (Bio-Rad). Image analysis was performed with PDQuest 2-D Analysis Software v8.0 (Bio-Rad). The molecular masses of proteins in gels were determined by co-electrophoresis of molecular weight standards (BenchMark Protein Ladder, Invitrogen, Carlsbad, CA, USA), while isoelectric point (pI) was determined by migration of protein spots on the linear IPG strips.

2.5. Liquid chromatography-tandem mass spectrometry analyses (LC-MS/MS)

Protein spots were carefully excised from the 2-DE gels and destained, reduced with 10 mM DTT in 25 mM ammonium bicarbonate followed by protein alkylation with 55 mM iodoacetamide. Protein digestion was carried out overnight at 37 °C with sequencing grade trypsin (Promega, Madison, WI, USA). Nanoscale LC separation of tryptic peptides was performed with a nanoACQUITY UPLC System (Waters, Milford, MA, USA) and LC-MS/MS analysis was carried out in a SYNAPT-HDMS Q-TOF (Waters) as previously reported (Huerta-

Ocampo et al., 2014) with brief modifications: Accurate mass data were collected in an alternating Data Dependent Acquisition mode (DDA). In low energy mode, data were collected at constant collision energy of 3 eV. In elevated-energy mode, the collision energy was ramped from 15 to 45 eV during 3 s of integration.

2.6. Protein identification

MS/MS spectra datasets were used to generate PKL files (.pkl) using ProteinLynx Global Server v2.4 (Waters). Proteins were then identified using PKL files and the MASCOT search engine v2.3 (Matrix Science, London, UK). Searches were conducted using the *Viridiplantae* subset of the NCBI protein database (2391213 sequences, December 2013) and an in-house database containing the *Jatropha curcas* nucleotide coding sequences available at the NCBI database (77340 sequences, December 2013). Trypsin was used as the specific protease and one missed cleavage was allowed. The mass tolerance for precursor and fragment ions was set to 100 ppm and 0.1 Da respectively and peptide charges were set at +2, +3, and +4. Carbamidomethylation of cysteine was set as fixed modification and oxidation of methionine was set as variable modification. Identifications were considered successful when significant MASCOT scores (> 33 for *J. curcas* nucleotide in-house and > 51 for *Viridiplantae* protein database) were obtained, indicating the identity or extensive homology at $p < 0.01$ and the presence of at least two peptides were considered necessary for reliable identification. Identified proteins were classified into different categories according to

Gene Ontology (<http://www.geneontology.org/>). A biological pathway diagram showing the identified enzymes involved in a metabolic network was generated using the visualization and analysis tool PathVisio v3.2.4 (Kutmon et al., 2015).

3. Results

3.1. Protein extraction

Although Osborne's classification based on solubility criteria is an ambiguous system, due that all depends in solutions used for extractions, is the most useful method for seed storage protein characterization and is a very useful method for protein pre-fractionation towards seeds storage proteins proteomics analysis (Jorriñ-Novó, 2014). Hence, the proteins in press-cake *J. curcas* were subjected to sequential extraction. It was observed that globulins fraction corresponds only to 20.17%, while glutelins represented the main fraction with 42.03% (Supplementary Table S1), which is different to all legumes where the main fraction is globulins. The representative electrophoretic profile in one dimension gel (SDS-PAGE) of different *J. curcas* press-cake protein fractions is shown in Fig. 1A.

3.2. 2-DE maps of seed protein fractions

The water-soluble protein fraction (albumins) was separated by 2-DE in the range of linear pH 5–8, while the salt soluble proteins (globulins) and glutelins were resolved in the range of linear pH 3–10. In the albumins fraction 434 spots were separated in 2-DE gels (Fig. 1B, Supplementary Fig. S2), while 310 spots for globulins (Fig. 1C, Supplementary Fig. S3) and 175 spots for glutelins (Fig. 1D, Supplementary Fig. S4) were resolved. From 84.1–96.0% of the protein spots from the three seed storage protein fractions were successfully identified by LC-MS/MS analysis and homology database search (Table 1). All identified proteins were classified into its functional classes. In albumins fraction were identified more than 100 protein species associated with well-known storage reservoir (Fig. 2A). Enzymes involved in glycolysis, TCA cycle. Enzymes related with sugars, nitrogen, and lipid metabolisms were also represented, as well proteins related with oxidoreduction function (SOD, GST). Proteasome subunits, 14-3-3 proteins, and enzymes related with defence and detoxification (LEA, glyoxalase, cyclophilin, and lactoylglutathione lyase) were also identified. Several heat shock proteins and proteases were identified (SERPIN-ZX-like, cysteine proteinase, leucine aminopeptidase). Interestingly in albumins fraction were detected enzymes related with the THF pathway and purine metabolism that have not been reported before (Table 2, Supplementary File S1).

Globulins and glutelins fractions showed 24 and 7 protein species, respectively (Fig. 2B and C) with several proteoforms representing each protein species. Globulins fraction was composed mainly by legumin B-like, 2S albumin-like and germin-like protein, curcin 1 and enzymes related to energy production such as ATP synthase and luminal-binding protein (Table 3 and Supplementary File S2). Glutelins, although the most abundant fraction in *J. curcas* press-cake was composed mainly of high molecular weight globulins (Table 4, Supplementary File S3). In Supplementary File S4 is shown the list of theoretical and experimental molecular weight/isoelectric point corresponding to all identified proteins.

Table 1

Protein identification of *Jatropha curcas* seed storage protein fractions (albumins, globulins and glutelins) by LC-MS/MS and homology database search.

Fraction	Spots	Identified (%)	Non-identified (%)	Unique
Albumins	434	365 (84.1)	69 (15.9)	100
Globulins	310	296 (95.5)	14 (4.5)	24
Glutelins	175	168 (96.0)	7 (5.0)	7

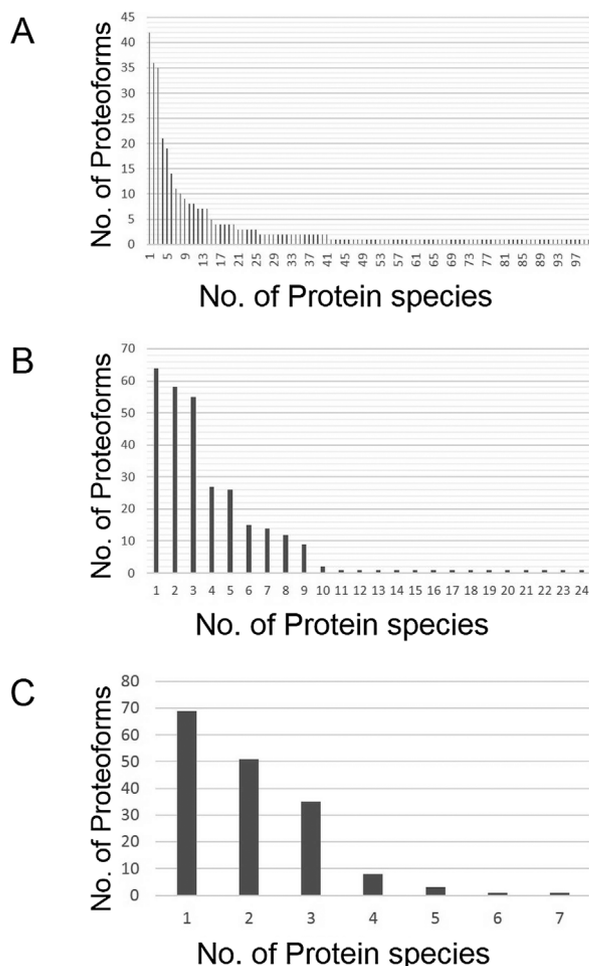


Fig. 2. Diagram indicating the number of proteoforms from different protein species detected in seed storage proteins. A) albumins, B) globulins, C) glutelins.

4. Discussion

4.1. *J. curcas* seeds protein fractionation

It is recognized that the total set of protein species and their corresponding proteoforms (frequently abundant) that constitute the cells proteome could not be captured in just one experiment (Romero-Rodríguez et al., 2014). Therefore, the use of fractionation techniques is necessary in order to obtain a more complete proteome characterization. Osborne's method has been long used for seed protein fractionation and is still one of the preferred procedures for protein pre-fractionation before proteomics profiling of seeds.

The high lipid content in seeds turns *J. curcas* into a high value crop, however the press-cake obtained from non-toxic *J. curcas* has the potential to be used as source of food. In this sense, seed protein storage proteins are the basis for seed protein characterization; hence the seed storage protein fractions in *J. curcas* non-toxic seeds were characterized. Most of enzymatic proteins identified were found in albumins fraction (water soluble-proteins). The network of metabolic activities of identified enzymes is represented in Fig. 3. Globulins and glutelins were represented by nutrient reservoir proteins (Table 3 and 4).

4.2. Protein species in albumins fraction

4.2.1. Protein species related to nutrient reservoir

Several proteoforms of legumin B and legumin A were detected (45 and 36, spots, respectively). 11S globulin was detected in 21 spots and globulin-1 S in 9 spots (Table 2). Globulin 1 (GLB1) is one of the most

Table 2
Identification by LC/MS–MS of albumins fraction proteins extracted from non-toxic *J. curcas* meal.

Protein name ^a	Spots ^b	<i>J. curcas</i> number ^c	NCBI accession number ^d	Theor. kDa/ pI ^e	Query Cover./E value ^f	Ident ^g
Nutrient Reservoir						
Legumin B-like precursor	42	Jcr4S00279.60	NP_001295688.1	57.5/6.75	100/0	100
Legumin B-like precursor	3	Jcr4S00279.80	NP_001295689.1	50.7/9.46	100/0	93
Legumin A-like	36	Jcr4S01636.60	XP_012085316.1	55.1/6.38	100/0	100
11S globulin seed storage protein 2-like	21	Jcr4S01636.40	XP_012085320.1	54.0/8.77	95/0	100
Globulin-1 S allele	2	Jcr4S03153.60	XP_012079069.1	78.2/5.38	66/0	100
Vicilin-like antimicrobial peptides 2-2	19	Jcr4S00603.20	XP_012064865.1	56.8/6.04	94/2 ^{e-128}	93
Vicilin-like antimicrobial peptides 2-2	1	Jcr4S03723.20	XP_012092200.1	50.7/6.04	100/0	86
2S albumin	35	Jcr4S00619.70	XP_012070746.1	16.2/5.53	100/2 ^{e-96}	100
2S albumin-like	1	Jcr4S00619.40	XP_012071216.1	16.2/8.38	100/2 ^{e-97}	100
2S albumin-like	1	Jcr4S00619.60	XP_012071217.1	16.6/6.81	100/1 ^{e-81}	100
2S albumin precursor, putative	1	Jcr4S00619.50	XP_012070745.1	16.4/6.88	100/1 ^{e-100}	100
Cupin 2	1	Jcr4S15278.20	XP_012079069.1	70.1/5.61	68/0	100
Em-like protein, protein SLE2	2	Jcr4S06850.20	XP_012087069.1	9.8/5.50	100/2 ^{e-60}	100
Curcin, 1Ribosome-inactivating protein cucurmosin-like precursor,	8	Jcr4S01069.20	NP_001295744.1	50.9/8.90	100/0	100
Glycolysis, TCA cycle						
Glyceraldehyde-3-P DH, cytosolic	14	Jcr4S00100.140	XP_012079890.1	44.5/7.05	82/9 ^{e-140}	99
Fructose-bisphosphate aldolase cytoplasmic	11	Jcr4S14120.10	XP_012066441.1	29.4/7.75	100/0	100
Enolase	8	Jcr4S00171.20	XP_012090397.1	52.3/5.71	100/0	100
Enolase-like	1	Jcr4S01799.10	XP_012090393.1	46.9/8.0	100/0	100
Enolase 1, chloroplastic	1	Jcr4S01892.10	XP_012092922.1	44.0/5.62	100/0	100
Triosephosphate isomerase, chloroplastic	1	Jcr4S02839.20	XP_012084383.1	41.6/7.60	100/0	100
Triosephosphate isomerase, cytosolic	1	Jcr4S19239.10	XP_012075954.1	36.5/8.07	100/0	100
2,3-bisphosphoglycerate-independent phosphoglycerate mutase	7	Jcr4S00101.20	XP_012090969.1	61.5/5.86	100/0	100
Phosphoglucomutase	1	Jcr4S02212.70	XP_012093262.1	32.8/6.23	94/0	97
Phosphoglucomutase, putative	2	Jcr4S15954.10	XP_012093260.1	35.9/8.84	89/2 ^{e-180}	100
Aldehyde dehydrogenase family 2 member B4, mitochondrial-like	2	Jcr4S03546.10	XP_012080093.1	59.5/8.26	94/0	100
Pyruvate dehydrogenase E1 component subunit beta-1, mitochondrial isoform X1	4	Jcr4S00168.90	XP_012088753.1	40.6/5.64	100/0	100
Cytosolic malate dehydrogenase	8	Jcr4S00279.20	XP_012069835.1	36.0/6.40	100/0	100
Malate dehydrogenase, mitochondrial	7	Jcr4S03295.10	XP_012078553.1	37.0/8.54	100/0	100
Aconitate hydratase, cytoplasmic	1	Jcr4S06787.10	XP_012089852.1	11.6/8.38	95/0	81
Citrate synthase, mitochondrial	1	Jcr4S00215.140	XP_012092602.1	49.4/6.33	100/0	100
Citrate synthase, mitochondrial	1	Jcr4S01687.10	XP_012092625.1	82.2/6.03	100/0	97
Carbohydrate metabolism						
UTP–glucose-1-phosphate uridylyltransferase	4	Jcr4S00684.10	XP_012074723.1	58.1/6.74	100/0	100
Sorbitol Dehydrogenase family protein	2	Jcr4S01523.120	XP_012090335.1	42.7/5.97	100/0	95
Beta-galactosidase 8	1	Jcr4S02941.10	XP_012064843.1	89.0/7.94	100/0	88
Beta-xylosidase/alpha-L-arabinofuranosidase 1	1	Jcr4S03968.30	XP_012080785.1	84.4/6.63	100/0	100
Aldose 1-epimerase-like	1	Jcr4S03178.30	XP_012086275.1	37.0/5.64	100/0	100
Lipid metabolism						
Acyl-CoA-binding protein, partial	1	Jcr4S00983.40	XP_012070513.1	9.4/10.0	81/3 ^{e-36}	88
Hydroxyacyl-ACP Dehydrase	1	Jcr4S03305.10	XP_012086368.1	9.0/8.93	100/3 ^{e-49}	100
Oil body-associated 2B-like (LOC105649726)	1	Jcr4S00805.80	XP_012091855.1	28.2/5.91	100/0	100
Amino acids and purine metabolism						
Ureidoglycolate hydrolase (LOC105635368)	2	Jcr4S03735.10	XP_012073838.1	21.9/5.54	100/6 ^{e-146}	100
Alanine aminotransferase 2-like	1	Jcr4S00131.20	XP_012068795.1	55.6/5.83	100/0	100
Nitrogen regulatory protein P-II	1	Jcr4S00680.140	XP_012082957.1	23.4/9.73	100/2 ^{e-130}	92
Reactive Intermediate Deaminase A, chloroplastic	1	Jcr4S00736.130	XP_012070823.1	22.2/8.99	100/3 ^{e-123}	91
Beta-ureidopropionase	1	Jcr4S00584.60	XP_012081364.1	49.3/5.86	100/0	100
Detoxification and plant resistance						
4-hydroxy-4-methyl-2-oxoglutarate aldolase 2 (putative)	1	Jcr4S00591.70	XP_012085762.1	17.9/5.68	100/1 ^{e-115}	99
Dienelactone hydrolase family protein	1	Jcr4S03008.10	XP_002321135.1	28.5/5.77	82/3 ^{e-124}	81
Lactoylglutathione lyase	1	Jcr4S01524.20	XP_012064721.1	26.0/8.50	100/5 ^{e-142}	100
Cyclophilin	2	Jcr4S01173.10	XP_012084373.1	18.3/8.68	100/4 ^{e-112}	100
Lactoylglutathione lyase, chloroplast isoform X1	2	Jcr4S00034.190	XP_012076751.1	34.2/8.91	100/0	100
Superoxide dismutase [Mn], mitochondrial-like	4	Jcr4S00073.110	NP_001295624.1	26.41/6.71	100/0	100
Cu/Zn superoxide dismutase	1	Jcr4S02704.30	NP_001295663.1	25.1/6.08	92/1 ^{e-130}	100
Cu/Zn superoxide dismutase family protein	1	Jcr4S00546.10	XP_012089157.1	20.7/6.74	100/4 ^{e-86}	77
Growth Regulation						
Thiamine thiazole synthase, chloroplastic	2	Jcr4S00619.120	XP_012070751.1	37.8/5.53	90/0	100
Phosphatidylethanolamine binding protein	1	Jcr4S05660.30	XP_012089808.1	18.4/5.14	100/5 ^{e-123}	100
Luminal-binding protein 5	2	Jcr4S22943.10	XP_012074649.1	73.6/5.11	96/0	100
Luminal-binding protein 5-like	1	Jcr4S00151.70	XP_012086670.1	69.9/5.14	100/0	100
Proteases						
Cysteine proteinase RD19a-like	1	Jcr4S00377.90	XP_012071959.1	41.5/5.81	100/0	100
Serpin-ZX-like	1	Jcr4S00079.140	XP_012071719.1	42.4/5.94	100/0	100
Leucine aminopeptidase 1-like	3	Jcr4S00343.100	XP_012071507.1	74.1/8.48	100/0	100
Oxide-Reduction						

(continued on next page)

Table 2 (continued)

Protein name ^a	Spots ^b	<i>J. curcas</i> number ^c	NCBI accession number ^d	Theor. kDa/ pI ^e	Query Cover./E value ^f	Ident ^g
Glutaredoxin GRX	10	Jcr4S06568.10	ADB02895.1	11.4/6.71	89/2e ⁻⁶⁴	100
Glutaredoxin	3	Jcr4S01536.70	XP_012080039.1	12.2/8.17	86/0	80
Thioredoxin H-type	1	Jcr4S04538.10	NP_001295646.1	21.1/8.54	68/5e ⁻⁶⁶	100
Thioredoxin reductase NTRB-like	1	Jcr4S09888.10	XP_012087579.1	35.3/9.35	97/2e ⁻¹⁶⁰	81
Protein disulfide-isomerase precursor	4	Jcr4S06094.20	NP_001295629.1	56.8/4.85	100/0	99
1-Cys peroxiredoxin	1	Jcr4S01479.40	XP_012067545.1	14.9/7.03	100/0	91
Peroxiredoxin	4	Jcr4S07807.20	XP_012067545.1	24.3/6.08	100/5e ⁻¹⁵⁸	100
Peroxidase 12-like	2	Jcr4S06050.30	XP_012073952.1	14.5/4.58	95/0	100
Ferritin, plant	1	Jcr4S03468.40	KDP22838.1	28.1/5.51	91/1e ⁻¹³²	100
Glutathione reductase, cytosolic	1	Jcr4S00410.70	XP_012071166.1	53.0/5.63	100/0	100
Glutathione S-transferase DHAR2-like	1	Jcr4S05806.20	NP_001295678.1	22.5/5.75	100/5e ⁻¹⁴⁶	100
Glutathione S-transferase parC (Hypothetical protein JCGZ_06274)	1	Jcr4S12402.20	KDP37218.1	25.7/6.02	100/2e ⁻¹⁶²	100
Glutathione-S-transferase theta, GST	1	Jcr4S03348.10	XP_012092523.1	24.3/6.24	100/2e ⁻¹⁵⁰	99
Glutathione-S-transferase theta, partial	1	Jcr4S00141.30	XP_012082716.1	24.2/6.24	100/2e ⁻¹³¹	98
Peroxiredoxin	1	Jcr4S00501.60	XP_012076732.1	17.6/5.15	100/2e ⁻⁹⁴	100
NAD(P)-dependent oxidoreductase (uncharacterized protein At5g02240)	1	Jcr4S05357.40	XP_012074957.1	23.9/5.94	100/0	100
Heat Shock						
Heat shock protein 83-like	7	Jcr4S00535.40	XP_012086831.1	78.9/4.98	98/0	100
Chaperonin CPN60-2, mitochondrial	4	Jcr4S02414.50	XP_012077667.1	58.0/5.9	100/0	91
FAM10 family protein At4g22670	3	Jcr4S00409.160	XP_012087117.1	52.9/5.21	100/0	100
Heat shock protein	2	Jcr4S08787.20	XP_012089450.1	65.2/5.26	100/0	91
Small heat shock protein	2	Jcr4S00024.10	ADT65203.1	15.3/7.79	84/3e ⁻⁵⁴	99
Small heat shock protein	2	Jcr4S00071.90	NP_001295605.1	58.5/7.06	58/8e ⁻¹⁰⁰	98
15.7 kDa heat shock protein, peroxisomal	1	Jcr4S02140.20	XP_012086420.1	15.9/6.85	100/7e ⁻⁹⁹	100
17.8 kDa class I heat shock protein-like	1	Jcr4S14146.30	XP_012088215.1	18.4/6.78	100/1e ⁻⁸⁷	100
Chaperonin	1	Jcr4S02314.20	XP_012077092.1	26.8/8.76	100/9e ⁻¹⁷⁶	100
Heat shock 70 kDa protein 15-like	1	Jcr4S00447.50	XP_012073334.1	92.7/5.24	100/0	100
Heat shock cognate 70 kDa protein 2-like	1	Jcr4S03314.10	XP_012089411.1	71.4/5.14	100/0	100
Nucleoside diphosphate kinase B	3	Jcr4S02985.40	NP_001295742.1	16.3/6.32	100/2e ⁻¹⁰⁴	100
Heat shock 70 kDa protein, mitochondrial	2	Jcr4S02598.30	XP_012074575.1	73.3/5.66	98/0	100
Ribosome, Proteasome						
Elongation factor 1-beta, putative	1	Jcr4S00111.50	XP_012079853.1	26.1/4.49	96/4e ⁻¹⁰²	100
Regulator of ribonuclease activity	1	Jcr4S00285.100	XP_012081861.1	17.7/5.33	100/7e ⁻¹¹⁵	100
Ribonuclease 3 family protein	1	Jcr4S01880.100	XP_012080098.1	17.5/9.75	86/1e ⁻⁷⁷	99
Proteasome subunit alpha type	1	Jcr4S01752.50	XP_012077064.1	27.4/5.92	100/0	100
Proteasome subunit alpha type 3 family protein	1	Jcr4S03336.20	XP_012075968.1	21.5/8.82	100/6e ⁻¹⁴²	99
Proteasome subunit alpha type-2A	1	Jcr4S02802.40	XP_012087375.1	24/5.30	100/1e ⁻¹⁷¹	100
Proteasome subunit alpha type-6-like	1	Jcr4S00843.120	XP_012079628.1	27.4/5.81	100/0	100
Proteasome subunit beta type	1	Jcr4S01070.40	XP_012079483.1	27.2/6.13	100/2e ⁻¹³⁷	82
Proteasome subunit beta type	1	Jcr4S00057.260	XP_012083566.1	24.9/6.95	100/1e ⁻¹⁵⁹	100
Small ubiquitin-related modifier 2-like	1	Jcr4U30955.10	XP_012080638.1	11.6/4.95	100/3e ⁻⁶⁹	100
14-3-3 protein 7	1	Jcr4S02291.60	XP_012067879.1	27.8/4.92	100/9e ⁻¹⁶⁹	99
Non characterized						
WD-40 like beta propeller (Uncharacterized protein LOC105638961)	2	Jcr4S00283.70	XP_012078271.1	77.6/6.15	100/0	100
DDPIV_N (uncharacterized protein LOC105642694 isoform X2)	1	Jcr4S01894.20	XP_012082992.1	56.1/5.61	100/0	97
Pentatricopeptide repeat-containing protein At4g14850	1	Jcr4S00033.30	XP_012069644.1	87.8/5.92	100/0	100

^aProtein names, ^bNumber of spots (Supplementary Fig. S2) where the same protein was identified (Supplementary File S1). ^cAccession numbers according to *Jatropha curcas* genome database (release 4.5, May 2014) were obtained after homology database identification for every single spot analyzed by LC-MS/MS. ^dNCBI accession numbers, ^eTheoretical mass (kDa) and pI of identified proteins, ^fQuery coverage, E-value and ^g% of identity were obtained after blastx alignment of *J. curcas* genome nucleotide sequences against de Viridiplante subset of the NCBI protein database.

abundant proteins accumulated in maize seed tissues and has been designated as an excellent indicator for the scrutiny of genetic variation (Hilton and Gaut, 1998). Two isoforms of vicilin-like antimicrobial peptides 2-2 (AMPs) were detected in 88 spots while 38 spots corresponded to 2S-albumins (Table 2). Antimicrobial peptides have been frequently isolated from seeds among other plant tissues (Franco et al., 2006; Lipkin et al., 2005; Pelegrini et al., 2006, 2008). It has been reported that *J. curcas* is a good source of cyclic peptides with diverse biological functions such as antimalarial, antiplatelet, antiproliferative, and cytotoxic (Pinto et al., 2015). Xiao et al. (2011) reported the JCpep7 peptide that displayed antimicrobial activity against Gram-negative pathogens like *Shigella dysenteriae*, *Pseudomonas aeruginosa* and *Salmonella typhimurium*. These bioactive peptides found in *J. curcas* could contribute in the development of antimicrobial strategies based in the use of novel natural sources (Mandal et al., 2009).

One LEA protein (Em-like) was detected in 2 spots. LEAs (Late embryogenesis abundant proteins) are widely recognized as associated

with desiccation tolerance in seeds. Shih et al. (2010) identified an Em-like protein in rice, which was not been identified previously in cereals, by other hand Vicent et al. (2000) reported two *Em* genes in Arabidopsis.

He et al. (2011) described that non-toxic *J. curcas* does not contain phorbol esters, nevertheless contains curcins. In agreement with this statement, some faint spots were detected in the *J. curcas* 2-DE map, which are not enough to cause the toxic effects. Lin et al. (2010) have indicated that curcin hemagglutinating activity is presented at more than 7.8 mg/L and LD₅₀ for oral semi-lethal dose was reported to be 104.73 mg/kg. Curcins type-1 are proteins that specifically are accumulated in the endosperm (Gu et al., 2015; Lin et al., 2003a) and are classified as Type I ribosome-inactivating proteins (RIP). Zhang et al. (2017) reported a novel RIP from *J. curcas*, however it is not well documented its function but it may be part of the response to biotic and abiotic stresses (Qin et al., 2009). Curcins have also related with anti-tumor activity (Lin et al., 2003b; Luo et al., 2006).

Table 3
Identification by LC-MS/MS of globulin fraction proteins extracted from non-toxic *Jatropha curcas* meal.

Protein name ^a	Spots ^b	<i>J. curcas</i> Number ^c	NCBIr Accession ^d	Theor kDa/pI ^e	Query coverage/ E-value ^f	Ident ^g
Nutrient reservoir						
Legumin B-like	64	Jcr4S00279.60	NP_001295688.1	57.5/6.75	100/0	100
Legumin B-like	27	Jcr4S00279.80	KDP40339.1	50.7/9.46	100/0	93
Legumin A-like	58	Jcr4S01636.60	XP_012085316.1	65.4/9.15	79/0	87
Legumin A-like	16	Jcr4S01636.70	XP_012085315.1	52.9/5.40	100/0	100
11S globulin seed storage protein	55	Jcr4S01636.40	XP_012085320.1	54.0/8.77	95/0	100
2S albumin-like	26	Jcr4S00619.70	XP_012070746.1	16.3/5.53	100/2e ⁻⁹⁶	100
2S albumin-like	10	Jcr4S00619.90	XP_012071219.1	20.4/8.30	98/9e ⁻⁶⁴	100
2S albumin-like	2	Jcr4S00619.60	XP_012071219.1	16.6/6.81	100/3e ⁻⁸⁰	100
2S albumin-like	1	Jcr4S00619.80	XP_012070747.1	16.6/6.34	85/5e ⁻⁸⁴	100
Vicilin-like antimicrobial peptides 2-2	14	Jcr4S00603.20	XP_012064865.1	57.0/6.04	100/0	100
Vicilin-like antimicrobial peptides 2-2	1	Jcr4S03723.20	XP_012092200.1	50.7/6.04	100/0	86
Germin-like protein 5-1	1	Jcr4S00027.200	XP_012088803.1	23.5/6.90	88/6e ⁻¹²²	99
Curcin 1	9	Jcr4S01069.20	ADN39429.1	51.0/8.90	60/0	100
Metabolism						
Glyceraldehyde-3-phosphate dehydrogenase, cytosolic	1	Jcr4S00100.140	XP_012079890.1	44.5/7.05	82/1e ⁻¹³⁹	99
Malate dehydrogenase	1	Jcr4S00279.20	XP_012069835.1	69.9/5.14	95/0	94
Malate dehydrogenase, mitochondrial	1	Jcr4S03295.10	XP_012078553.1	37.0/8.54	96/0	99
Oxide-reduction						
Protein disulfide-isomerase precursor	2	Jcr4S06094.20	NP_001295629.1	56.9/4.85	100/0	100
Superoxide dismutase [Mn], mitochondrial-like	1	Jcr4S00073.110	NP_001295624.1	30.4/6.79	100/1e ⁻¹⁶³	87
Glutaredoxin GRX	1	Jcr4S06568.10	NP_001295635.1	11.47/6.71	89/2e ⁻⁶⁴	100
Ribosomal, energy						
60S acidic ribosomal protein P3-like	1	Jcr4S16124.30	XP_012075150.1	12.2/4.24	100/1e ⁻²⁹	100
ATP synthase subunit beta, mitochondrial	1	Jcr4S01269.80	XP_012075690.1	61.1/6.30	89/0	96
ATP synthase subunit beta, mitochondrial-like	1	Jcr4S00914.20	XP_012066038.1	59.7/6.06	94/0	100
AtpB (chloroplast)	1	Jcr4S00559.20	YP_002720119.1	50.0/4.91	97/0	99
Luminal-binding protein 5	1	Jcr4S22943.10	XP_012074649.1	73.7/5.11	100/0	100
Luminal-binding protein 5-like	1	Jcr4S00151.70	XP_012086670.1	69.9/5.14	95/0	94
Non identified	13	–	–	–	–	–

^aProtein names, ^bNumber of spots (Supplementary Fig. S3) where the same protein was identified (Supplementary File S2). ^cAccession numbers according to *Jatropha curcas* genome database (release 4.5, May 2014) were obtained after homology database identification for every single spot analyzed by LC-MS/MS. ^dNCBIr accession numbers, ^eTheoretical mass (kDa) and pI of identified proteins, ^fQuery coverage, E-value and ^gPercentage of identity were obtained after blastx alignment of *J. curcas* genome nucleotide sequences against de Viridiplante subset of the NCBIr protein database.

4.2.2. Protein species involved in sugar metabolism

Glycolysis is well represented in *J. curcas* albumins fraction, glyceraldehyde-3P-DH was detected in 14 spots, two isoforms of phosphoglucosmutase and triose phosphate isomerase and three isoforms of enolase were identified. The pyruvate dehydrogenase E1 component subunit beta-1, part of the complex that carried out the conversion of pyruvate to Acetyl-CoA, was detected in four spots. Aldehyde dehydrogenase, phosphoglycerate mutase, and enzymes related to the TCA cycle (aconitase, two isoforms of citrate synthase and two isoforms of malate dehydrogenase) were identified (Table 2).

Uridyltransferase (UTP-glucose-1-phosphate uridylyltransferase) or glucose-1-phosphate uridylyltransferase (detected in 2 spots) is an enzyme involved in carbohydrate metabolism with an important role in glycogenesis and cell wall synthesis (Dai et al., 2006). In relation to the

sugars metabolism, beta-galactosidase 8 and beta-xylosidase/alpha-L-arabinofuranosidase 1 were detected in *J. curcas*. Aldose-1-epimerase, which catalyses the first step in galactose metabolism was also detected in one spot. Two proteoforms of sorbitol dehydrogenase enzyme, which converts sorbitol into fructose, were also detected.

4.2.3. Protein species related to fatty acid metabolism

Proteins related to fatty acid metabolism were identified in albumins fraction (Table 2). Among them, the hydroxyacyl-ACP-dehydrase, is the key enzyme of the fatty acid synthesis (FAS) system (Lung and Chye, 2016) was identified in one spot. One proteoform of the acyl-Co-A-binding protein was identified in *J. curcas* albumins fraction (Table 2), this is an enzyme that binds medium- and long- chain acyl CoA esters with very high affinity. Acyl-Co-A-binding protein may

Table 4
Identification by LC-MS/MS of glutelin fraction proteins extracted from non-toxic *J. curcas* meal.

Protein name ^a	Spots ^b	<i>J. curcas</i> number ^c	NCBI accession number ^d	Theor. kDa/pI ^e	Query coverage/E value ^f	Ident ^g
Legumin B-like precursor	69	Jcr4S00279.60	NP_001295688.1	57.5/6.75	100/0	99
Vicilin-like antimicrobial peptides 2-2	3	Jcr4S00603.20	XP_012064865.1	56.8/6.04	100/0	100
2S albumin-like	1	Jcr4S00619.70	XP_012070746.1	16.2/5.53	100/2e ⁻⁹⁶	100
Legumin A-like	8	Jcr4S01636.70	XP_012085315.1	52.9/5.40	100/0	100
Legumin A-like	51	Jcr4S01636.60	XP_012085316.1	65.4/9.15	79/0	87
11S globulin seed storage protein 2-like	35	Jcr4S01636.40	XP_012085320.1	54.0/8.77	95/0	100
Not identified	7	–	–	–	–	–

^aProtein names, ^bNumber of spots (Supplementary Fig. S4) where the same protein was identified (Supplementary File S3). ^cAccession numbers according to *Jatropha curcas* genome database (release 4.5, May 2014) were obtained after homology database identification for every single spot analyzed by LC-MS/MS. ^dNCBIr accession numbers, ^eTheoretical mass (kDa) and pI of identified proteins, ^fQuery coverage, E-value and ^gPercentage of identity were obtained after blastx alignment of *J. curcas* genome nucleotide sequences against de Viridiplante subset of the NCBIr protein database.

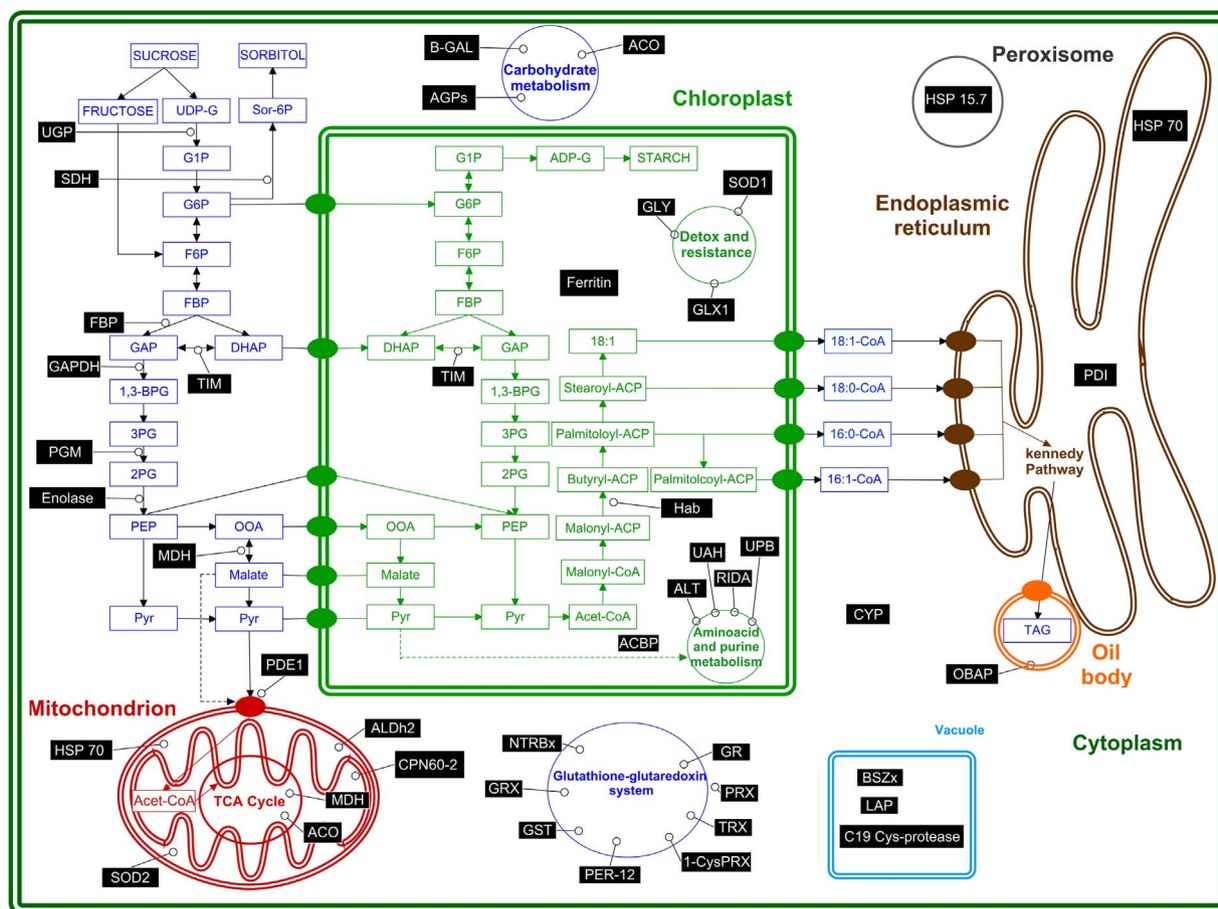


Fig. 3. Metabolic network of non-toxic *Jatropha curcas* seeds. The identified protein spots are indicated in black spots (Table 2). Diagram was generated using the visualization and analysis tool PathVisio 3.2.4.

function as maintenance, protection, and transport of the Acyl-CoA pool (Yurchenko and Weselake, 2011).

In *J. curcas* seeds the triacylglycerol's (TAGs) are accumulated in oil bodies, which consist of a central core of neutral lipids, delimited by a monolayer of phospho- and glyco-lipids including sterols and proteins associated to the oil bodies surface. One proteoform of the oil body-associated 2B-like protein in albumins fraction was identified. In maize (*Zea mays* L.), the oil body associated protein 1 gene (*obap1*) was found mainly expressed in scutellum during maturation (López-Ribera et al., 2014).

4.2.4. Protein species related to amino acids and purine metabolism

Nitrogen in seeds does not only come from root nitrogen uptake during the later growing period, but also from the redistribution from the vegetative organs, in addition, the recycling and redistribution of nitrogen in plants is important for the environmental stress response (Zhang et al., 2017). Four proteoforms related to amino acids and purine metabolism were identified in albumins fraction: Ureidoglycolate hydrolase, Alanine aminotransferase 2-like, Nitrogen regulatory protein P-II, and Reactive Intermediate Deaminase A, chloroplastic. Ureidoglycolate (two proteoforms) is an intermediate of purine catabolism catalyses the final step of ureide degradation in which inorganic nitrogen is re-assimilated (Li et al., 2015; Werner et al., 2013). The beta-propionase enzyme, detected in *J. curcas* albumins fraction (one spot), acts on carbon-nitrogen bonds, specifically in linear amides and participates in pyrimidine, beta-alanine metabolism, and pantothenate biosynthesis (Shen et al., 2014).

Alanine amino transferase, an enzyme belonging to the L-alanine degradation and the nitrogen regulatory protein P-II, a constituent of the adenylation cascade involved in the regulation of glutamine

synthetase (GS) activity were detected in one isoform (Table 2). In nitrogen-limiting conditions, P-II is uridylylated establishing a complex that allows the deadenylation of GS, thus activating the enzyme. On the contrary, under nitrogen surplus, this complex is deuridylylated promoting adenylation and the consequent inactivation of GS (Brown et al., 1971; Jonsson and Nordlund, 2007). In plants, the function of reactive intermediate deaminase A is still unclear, however recent reports allowed to established its contribution in the biosynthesis of branched-chain amino acids (Niehaus et al., 2014), one proteoform was detected in *J. curcas* press-cake albumins fraction.

4.2.5. Protein species involved in detoxification and plant resistance

4-hydroxy-4-methyl-2-oxoglutarate aldolase (4HMO), dienelactone hydrolase, lactoylglutathione lyase, and cyclophilin were detected in *J. curcas* albumins fraction. The enzyme 4HMO cleaves 4-carboxy-4-hydroxy-2-oxoglutarate giving as a product pyruvate. Participates in benzoate degradation via hydroxylation, which links aromatic catabolism to central cellular metabolism (Tack et al., 1972). Dienelactone hydrolases play a crucial role in the bacterial degradation of chloroaromatic compounds. Many representatives of this group of xenobiotic compounds are converted to chlorosubstituted catechols by the initial enzymes of aromatic catabolism (Schlöman et al., 1990). Two isoforms of lactoylglutathione lyase were detected, this is an enzyme that catalyses the isomerization of hemithioacetal adducts, which are cytotoxic compounds generated under abiotic stresses, including metal toxicity (Nahar et al., 2017). Cyclophilin (two isoforms) belongs to the immunophilin superfamily with peptidyl-prolyl *cis-trans* isomerase (PPIase activity). Catalyse the interconversion of the *cis-* and *trans-* rotamers of the peptidyl-prolyl amide bond of peptides. The interaction of soybean cyclophilin GmCYP1 with the isoflavonoid regulator

GmMYB176 and 14-3-3 proteins suggests its role in defence in soybean (Mainali et al., 2017). Interestingly, 14-3-3 proteins were also detected in *J. curcas*.

4.2.6. Protein species related to plant growth regulation

Thiamine thiazole synthase (TH1, two proteoforms) plays a central in thiamine biosynthesis, but it is also important in abiotic stress responses and mitochondrial DNA damage tolerance (Li et al., 2016). It has also been associated with heat tolerance in rice (Chen et al., 2011) and is involved in ABA-regulated stomatal movement and in the plants drought response (Li et al., 2016). Phosphatidylethanolamine binding protein (one isoform) includes various functions such as lipid binding, neuronal development, control of the switch between shoot growth and flower structures and the regulation of signalling such as the MPA kinase and the NF-kappaB pathways (Vallée et al., 2003; Yeung et al., 2001). Luminal-binding protein detected in three proteoforms, is involved in storage of calcium pool inside the endoplasmic reticulum lumen, accumulation of this protein confers resistance to drought (Valente et al., 2009). In soybean root tips the induction of the luminal-binding protein 5 was observed during flood conditions (Komatsu et al., 2012).

4.2.7. Proteases and oxidoreductases

Cysteine protease RD19a-like, leucine aminopeptidase (three proteoforms), and serine proteinase inhibitor (SERPIN-ZX-like) were detected in *J. curcas* albumins fraction. Cysteine proteases RD19a-like (one proteoform) or cathepsin F-like proteases are members of the Papain-like cysteine protease, a large class of proteolytic enzymes associated with several plant processes (Richau et al., 2012). Serine protease inhibitor (Serpin, one proteoform) is a family of proteases inhibitors widely distributed. Members of the group of leucine aminopeptidase (three proteoforms), are implicated in transcriptional regulation (Asqui et al., 2017).

Glutathione-S-transferases (GSTs), detected in *J. curcas* albumins (four proteoforms), are a family of phase II detoxification enzymes. The acquired resistance to chemotherapy, herbicides, insecticides and microbial antibiotics has been attributed to the presence of GST (Townsend and Tew, 2003). Glutaredoxin (13 isoforms) have similar functions of thioredoxin (TRX), both are proteins involved in cell protection against oxidative stress damage (Sánchez-Riego et al., 2016). Protein disulphide isomerase (four proteoforms) participates throughout the maturation of extracellular proteins adding disulphide bonds to stabilize it or to covalently join it to other proteins (Wang and Tsou, 1993). Glutathione reductase (GR, spot 23) or glutathione-disulphide reductase (GSR) catalyses the reduction of glutathione disulphide (GSSG) to the sulfhydryl form glutathione (GSH), which is a critical molecule in resisting oxidative stress and the maintenance of the cell reduced environment (Lüersen et al., 2013).

Iron is an essential nutrient for all cells, but its excess is harmful to cells, and so iron homeostasis must be controlled. Ferritins play important roles in sequestering or releasing iron as needed (Borg et al., 2012) and are exclusively targeted to plastids and mitochondria (Borg et al., 2012; Briat et al., 2010; Zancani et al., 2004). In this work, a ferritin was identified in albumins fraction (one proteoform). A recent report indicates that ferritins from wheat (*Triticum aestivum* L.) play important roles in enhancing tolerance in stresses associated with ROS (Zang et al., 2017).

4.2.8. Heat shock protein, ribosome, energy, and proteasome

A set of several proteoforms of heat shock proteins, small heat shock proteins, and chaperonins were identified in albumins fraction (Table 2). The nucleoside diphosphate kinase B (NDPKs, three isoforms) is an enzyme that catalyses, in a reversible manner, the production of different nucleoside diphosphates (NDP) and triphosphates (NTP) (Kihara et al., 2011). The regulator of ribonuclease (one proteoform) activity acts as a regulator of the endonuclease RNase E and inhibiting

the RNA processing. Ribonuclease 3 is a family of ribonucleases that recognizes dsRNA and cleaves it to transform them into mature RNAs, this enzyme was detected in one proteoform. Ubiquitin-fold modifier conjugating (UBL, one isoform) is covalently linked to target proteins but its physiological functions are still not known (Daniel and Liebau, 2014).

4.3. Protein species in globulins and glutelins fraction

Globulins fractions were represented by legumin A and B, 2S albumins, vicilin-like antimicrobial peptides, Germin-like protein 5-1, and curcin 1. Germins (GER) and germin-like proteins (GLPs), together with sucrose-binding proteins and seed globulins, are part of protein superfamily called cupin. Large numbers of GERs and GLPs have been functionally characterized from diverse plant species, their accumulation is related to biotic and abiotic stress response (Ilyas et al., 2016). Despite their solubility properties, globulins and glutelins are structurally similar, even evidences indicate that both protein share a common gene origin (Okita et al., 1989). Our results indicate that similar proteins were identified in both fractions (Table 3 and 4).

5. Conclusions

In the present study, the protein composition of seed storage proteins of non-toxic *J. curcas* press-cake after oil extraction was investigated using proteomics tools. Results have shown that albumins or water-soluble proteins are rich on different protein species related with carbohydrate metabolism. Other proteins involved in lipid, nitrogen, and purine metabolism were also identified. Proteins related with detoxification of xenobiotics were detected. Interestingly, the 14-3-3 proteins and NDPK, which are components of signaling pathways as well as cyclophilin that is one of protein targets of 14-3-3 were detected in *J. curcas* water soluble fraction. Although toxic protein such as curcin was detected, the amount present in the non-toxic *J. curcas* seeds may be not enough to cause toxic effects. This study contributes to the information of *J. curcas* seed proteome.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <https://doi.org/10.1016/j.indcrop.2017.11.046>.

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