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Abstract:	Helianthinins are storage proteins present in Helianthus annuus seeds belonging to the 11S globulin family. We describe here that a fraction of the helianthinins is phosphorylated. This conclusion is supported by different criteria including identification by MALDI-TOF of major protein bands revealed with a specific dye for phosphoproteins, anti-phosphoserine antibody and binding to a phosphoprotein affinity matrix. Moreover, we show that the phosphorylation status of helianthinins changes following germination.				

*Response to Referee/Editor Comments

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Manuscript number: SSR-D-13-00031R1

Title: Phosphorylated 11S globulins in sunflower seeds

Editor's decision: Minor Revision

Dear Dr. Derek Bewley,

We appreciate the reconsideration of the MS entitled Phosphorylated 11S globulins in

sunflower seeds, by Quiroga et al.

Reviewers requested a few minor revisions that have been taken into account and modified

in the present version. We hope that the changes introduced will fully meet the requirements

of SSR.

Kind regards,

Mariana Regente

Response to the Reviewers

Comments to Author:

Thank you for making the changes as requested. There are still a few minor revisions that

need to be made, particularly in the Tables and Figure legends, but then the paper is

acceptable.

Answer: All the suggestions of the reviewer have been taken into account and modified in

the revised MS.

Phosphorylated 11S globulins in sunflower seeds Running head title: Phosphorylation of sunflower storage proteins. Ivana Quiroga^{†a}, Mariana Regente^{†a}, Luciana Pagnussat^a, Ana Maldonado^b, Jesús Jorrín^b, Laura de la Canal^a*. ^a Instituto de Investigaciones Biológicas, Universidad Nacional de Mar del Plata-CONICET, Mar del Plata, Argentina. ^b Departamento de Bioquímica y Biología Molecular, Universidad de Córdoba, Córdoba, España. † Both authors have equally contributed to this work * Corresponding author Mariana Regente. Instituto de Investigaciones Biológicas, Universidad Nacional de Mar del Plata -CONICET, Funes 3250, 7600 Mar del Plata, Argentina. Tel 54-223-4753030. Fax 54-223-4724143. E-mail address: mregente@mdp.edu.ar **Keywords** Helianthus annuus, Germination, 11S Globulins, Helianthinin, Phosphorylation, Storage proteins.

Abstract

Phosphorylation of sunflower storage proteins.

Helianthinins are storage proteins present in *Helianthus annuus* seeds belonging to the 11S globulin family. We describe here that a fraction of the helianthinins is phosphorylated. This conclusion is supported by different criteria including identification by MALDI-TOF of major protein bands revealed with a specific dye for phosphoproteins, anti-phosphoserine antibody and binding to a phosphoprotein affinity matrix. Moreover, we show that the phosphorylation status of helianthinins changes following germination.

Introduction

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Seed storage proteins accumulate in high amounts during the late stages of seed development and provide nitrogen for seed germination and early seedling growth. These proteins have been classified into albumins, globulins and glutelins based on their solubility in water, saline, and NaOH solutions respectively (Osborne, 1924). Helianthinin, the main storage protein of sunflower seeds, belongs to the salt soluble 11S globulin fraction (Raymond et al., 1995). These globulins are composed by α and β subunits assembled as a hexamer (Shewry et al., 1995). They are synthesized as a precursor in the rough endoplasmic reticulum and follow the secretory pathway to be delivered into storage vacuoles. Sequence analysis revealed that the gene encoding the sunflower 11 S precursor (Helianthinin G3) contains a signal peptide responsible for its targeting to the secretory pathway, followed by the coding sequence for α and β chains (Vonder Haar et al., 1988). In fact, the proglobulin containing the α and β polypeptides is proteolytically cleaved after disulfide bond formation and transported through the secretory system as intermediate trimers that further assemble into the mature hexameric form (Shewry et al., 1995). On the other hand, helianthinins are polymorphic and their subunit composition varies among different sunflower cultivars. At least six types of subunits with molecular masses ranging from 40 to 64 kDa have been described so far, consisting in different combinations of the large (32-40 kDa) acidic α polypeptide and the small (20-25 kDa) basic β polypeptide (Rahma and Narasinga, 1979; Schwenke et al., 1979). Reversible protein phosphorylation is one of the most frequent post-translational modifications and its relevance is that it regulates diverse cellular processes. Protein phosphorylation in eukaryotes mostly occurs by the addition of a phosphate group on certain serine and threonine residues, whereas phosphorylation on tyrosine residues is less abundant. Post-translational modifications of a protein can determine its activity state, localization, turnover, and interactions with other proteins (Zolnierowics and Bollen, 2000). Besides its well-known role in several signaling pathways activating and deactivating enzymes, phosphorylation participates in an expanding catalogue of physiological functions in plants (Kline-Jonakin et al., 2011). Among them, it plays a key role during embryogenesis and seed germination (Fujii and Zhu, 2009; Nakashima et al., 2009). Also, Wan et al. (2007) have shown that the 12S globulin cruciferin is the major phosphorylated protein in Arabidopsis thaliana seeds. A total of 20 phosphorylation sites were identified on this globulin, including serine, threonine and tyrosine residues (Wan et al., 2007).

1 Although a specific function was not assigned to this post-translational modification, cruciferin 2 processing and mobilization were suggested. In this respect, Ghelis et al. (2008) detected changes 3 in cruciferin tyrosine-phosphorylation in response to ABA treatment, suggesting this is induced by 4 ABA to prevent cruciferin proteolysis. Agrawal and Thelen (2006) identified in a proteomic study 5 of B. napus phosphoproteins belonging to 10 major functional categories, including storage 6 proteins during seed filling. A recent analysis of seed maturation in Arabidopsis, rapeseed, and 7 soybean detected novel orthologs of seed storage proteins using a phosphoproteomic approach 8 (Meyer et al., 2012). 9 Taking into account these observations, the aim of this work was to elucidate whether

phosphorylated storage proteins are present in sunflower, a phylogenetically distant species.

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Materials and Methods

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Protein sample preparation and analysis

- 4 Helianthus annuus L. seeds (line 10347 Advanta Semillas SAIC) were sterilized in 27.5 mg/ml
- 5 sodium hypochlorite for 30 min, rinsed with sterile water and subjected to imbibition for the
- 6 indicated times at 25° C. Petri dishes containing wet Whatman filter paper under sterile conditions
- 7 were used. Dried seeds, germinating seeds (16 and 24 h of imbibition) and seedlings (48, 90 and
- 8 96 h of imbibition) were subjected to pulverization and extracted in the following buffer: 50 mM
- 9 Tris-HCl pH 7.5, 2 mM DTT, 0.1 M EDTA, protease and phosphatase inhibitors (Martin et al.,
- 10 2007).
- 11 Protein concentration was determined by the bicinchoninic acid method (Smith et al., 1985) using
- 12 bovine serum albumin as standard. Protein fractions (50 μg) were resolved in 12 % SDS-PAGE
- 13 (Laemmli, 1970) and stained with Coomassie Brilliant Blue R-250 or Colloidal Coomassie Blue
- 14 (Neuhoff *et al.*, 1988). Western blot assays were carried out by protein transfer onto nitrocellulose
- 15 membranes. Rabbit anti-phosphoserine 1:500 (Invitrogen) or sunflower anti-11S globulin
- antibodies provided by Dr. S. Petruccelli (1:1000) were used as primary antibodies and an anti-
- 17 rabbit IgG coupled to alkaline phosphatase (Sigma) was used as secondary antibody (1:10000).

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2D-PAGE

- 20 Homogenates from imbibed seeds (100 µg protein) were subjected to chloroform-methanol
- 21 precipitation and the pellet was suspended in 7 M urea, 2 M thiourea, 4 % (w/v) CHAPS, 100 mM
- 22 DTT, 0.2 % (v/v) ampholytes pH 3-10 (BioRad), trace of bromophenol blue. Samples were loaded
- onto pH 3–10 nonlinear, immobilized pH gradient strips (7 cm, ReadyStrips, BioRad). Isoelectric
- 24 focusing was performed applying the following conditions: for the rehydration step the voltage
- was maintained for 16 h at 50 V, and then the proteins were focused at 9000 V/h at 20°C. After
- isoelectric focusing the strips were equilibrated in 375 mM Tris-HCl pH 8.8, 6 M urea, 20 % (v/v)
- 27 glycerol, 2 % (w/v) SDS, 2 % (w/v) DTT, followed by 375 mM Tris-HCl pH 8.8, 6 M urea, 20 %
- 28 (v/v) glycerol, 2 % (w/v) SDS, 2.5 % (w/v) iodoacetamide. Second dimension was carried out
- 29 according to Laemmli (1970) in 12 % gels at 150 V and PeppermintStick phosphoprotein
- 30 molecular mass standards (Invitrogen) were loaded in the first gel lane. After SDS-PAGE, gels
- were fixed with 50 % (v/v) methanol and 10 % (v/v) acetic acid for 30 min, washed with distilled
- 32 water, stained with Pro Q Diamond phosphoprotein stain (Life Technologies) according to the

- 1 manufacturer's instructions and scanned with a FluorImager instrument (Amersham Biosciences)
- 2 using 514 nm laser as excitation source and a 570 nm band-pass emission filter. After, gels were
- 3 stained with SYPRO Ruby (Life Technologies) and scanned using a 488 laser as excitation source
- 4 and a 610 band-pass emission filter.

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MALDI-TOF

- 7 Protein bands and spots indicated in the text were automatically excised from 1D and 2D gels
- 8 employing Investigator ProPic robotic workstation (Genomic Solutions, UK). Gel pieces were
- 9 digested with trypsin according to standard protocols in a ProGest station (Genomic Solutions).
- 10 MS analyses of peptides were performed in a 4700 Proteomics Station (Applied Biosystems,
- USA) as previously described (Pérez-Reinado et al., 2007). Protein identification was assigned by
- comparing the obtained peptide mass fingerprinting with the non-redundant plant database, using a
- 13 Mascot 1.9 search engine (Matrixscience, UK) (Pinedo *et al.*, 2012).

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Phosphoprotein purification

- 16 Phosphoprotein enrichment was performed employing phosphoprotein affinity columns (Qiagen)
- according to the manufacturer's instructions and the previously described modifications adapted
- for plant extracts (Meimoun et al., 2007). Seeds were decorticated and protein was extracted using
- 19 25 mM MES, 1 M NaCl, 0.25 % (w/v) CHAPS, protease inhibitors and Benzonase Nuclease stock
- 20 solution (Phosphoprotein Purification Kit, Qiagen). Protein extracts (5 mg) were loaded onto a
- 21 Qiagen column and after washing with loading buffer (8 ml) bound phosphoproteins were eluted
- with 3 ml of elution buffer (Qiagen) and submitted to MALDI TOF for identification.

Results and Discussion

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3 As a first approach to detect putative phosphorylated proteins in sunflower seeds we performed 2D 4 SDS-PAGE of 16-h-imbibed seed protein extracts and the resulting gels were sequentially stained 5 for phosphoprotein detection (Pro-Q Diamond stain) and total protein (Sypro Ruby). Pro-Q 6 Diamond phosphoprotein stain allows in-gel detection of phosphate groups attached to tyrosine, 7 serine or threonine residues (Oh et al., 2009; Chitteti and Peng, 2007; Laugesen et al., 2006). Fig. 8 1B shows that a limited number of spots were clearly detected upon Pro-Q Diamond staining. 9 Among them, three spots that were consistently reproducible and could be major seed proteins 10 were further analyzed by peptide mass fingerprinting to identify them. They were digested with 11 trypsin, submitted to MALDI-TOF and putatively identified using Mascot software on the 12 Genbank database. Table 1 summarizes the data obtained. The three spots matched to the 11S 13 globulin seed storage protein G3 (Helianthinin G3) from *Helianthus annuus* (Uniprot ID P19084). 14 Also, good agreement was observed between the experimentally-determined molecular mass of 15 those spots and that predicted for β subunit of Helianthinin G3 (20.98 kDa). Helianthinin G3 16 corresponds to the precursor protein of 493 amino acids which, as described for other 11S 17 globulins, is further cleaved into two chains: the acidic chain (α subunit), and the basic chain (β 18 subunit) (Rahma and Narasinga, 1979; Schwenke et al., 1979; Vonder Haar et al., 1988; Shewry et 19 al., 1995). Indeed, according to proteomic criteria all the three spots detected with the 20 phosphoprotein specific stain can be assigned as 11S globulins. The phosphorylated spots between 21 35-40 kDa correlate with major spots in Fig 1A (total proteins). One major spot (~35 kDa and acid 22 isoelectric point) was clearly visible under Pro-Q Diamond staining, whose molecular mass and 23 isoelectric point are consistent with the \alpha subunit. Even though these spots could not be identified 24 by peptide mass fingerprinting, their features suggest that they correspond to α subunits of 11S 25 globulins. 26 Supporting the suggestion that 11S globulin subunits are phosphorylated in vivo, a bioinformatic 27 tool (NetPhos program, http://www.cbs.dtu.dk/services/NetPhos/) (Blom et al., 1999) predicts 16 28 putative phosphorylation sites on the Helianthinin G3 precursor sequence. These correspond to 14 29 serines, 1 threonine and 1 tyrosine (Fig. 2), covering both α and β subunits. To experimentally 30 confirm this presumption we used a strategy based on the purification of phosphoproteins by 31 affinity chromatography; the robustness of this procedure has been reported (Laugesen et al., 2006 32 Meimoun et al., 2007). Total seed extracts were loaded onto a Qiagen affinity matrix and eluted proteins were fractionated by SDS-PAGE followed by MALDI-TOF analysis of the major proteins. Among them, four bands could be identified and all matched to the 11S globulin precursor G3 (Table 2). Thus several criteria indicate that 11S globulins from imbibed sunflower seeds are phosphorylated: they are detected by a phosphoprotein specific stain, they are retained in a specific affinity matrix and, in both cases, their identity can be confirmed by peptide mass fingerprinting.

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In Arabidopsis the phosphorylation of cruciferins involves key residues allowing protein assembly, processing and mobilization (Wan et al., 2007), and changes in phosphorylation are regulated by ABA treatment (Ghelis et al., 2008). In this context, it can be hypothesized that globulins can vary in their phosphorylation status according to their physiological state. Thus, we have investigated whether phosphorylation of helianthinins can be regulated during seed germination and seedling growth. We have compared the total seed protein profile (Fig. 3A) and that of phosphoproteins detected with an anti-P-Ser antibody (Fig. 3B). As observed in Fig 3A, the expected protein pattern was observed in dry seeds (Raymond et al., 1995; Molina et al., 2004; Serre et al., 2001) with major bands of around 32-40 and 20-25 kDa corresponding to 11S globulin α and β subunits, respectively. In addition, there were bands of around 60 kDa that could correspond to an αβ subunit (Molina et al., 2004). The identity of these bands was also confirmed by immunoblotting because they were recognized by sunflower anti-11S globulins serum (not shown). The protein pattern in dry seeds does not change during the first 24 h of imbibition, but modification in the protein profile occurs thereafter (Fig. 3 A). Particularly, bands around 40 kDa (α subunit) consistently decrease due to their mobilization, as previously reported (Shewry et al., 1995). The 11S subunits appear to be differently phosphorylated during seedling growth while the band of around 60 kDa retains the same phosphorylation status (Fig. 3B). Even if bands of ~22 kDa (β subunit) are barely detected by the anti-P-Ser antibodies, subunits around 40-kDa are clearly visible from 16 to 48 h of seed imbibition while they are almost undetectable in dried seeds and after 90-96 h. The absence of phosphorylated α subunits after 2 days could be attributed to their degradation, since they are also undetectable by Coomassie Blue staining (Fig 3A). Dry seeds show a typical pattern of helianthinins but are barely detected by anti-P-Ser.

In conclusion, the phosphorylation of helianthinins appears to be induced following the completion of germination. Even though the basis and role of this post-translational modification is unknown, our results are indicative of the necessity for future investigations to analyze the

- 1 relationship between storage protein phosphorylation and mobilization during post-germinative
- 2 growth.

1 Acknowledgements

- 2 The authors thank Dr. Silvana Petruccelli for providing the sunflower anti-11S globulins antiboby.
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- 5 the University of Mar del Plata (Argentina).

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Figure legends Figure 1 2D gel electrophoresis of protein extracts from 16-h-imbibed sunflower seeds. (A) Sypro Ruby staining, (B) Pro-Q diamond staining. Location of phosphorylated molecular mass standards in kDa is indicated on the left. Circles indicate the spots identified by MALDI-TOF peptide mass fingerprinting. Figure 2 Amino acid sequence of 11S globulin seed storage protein G3 from Helianthus annuus (UniProt accession number P19084). Putative phosphorylated residues were predicted using the NetPhos 2.0 program and appear underlined in the sequence. (S: 14, T: 1, Y: 1). The signal peptide is shown in dark gray boxes; α and β subunits are presented in gray and white boxes respectively. Figure 3 Phosphoserine immunodetection of sunflower seed proteins at different times of germination and seedling growth. (A) 50 µg of total proteins extracted from dry seeds and after 16, 24, 48, 90 and 96 h of imbibition were fractionated by SDS-PAGE and stained with Coomassie Blue. (B) Immunoblotting with anti-phosphoserine antibody of the same samples detailed in (A). Figure shows a representative experiment of three replicates. Molecular mass in kDa is indicated on the left.

Figure 1

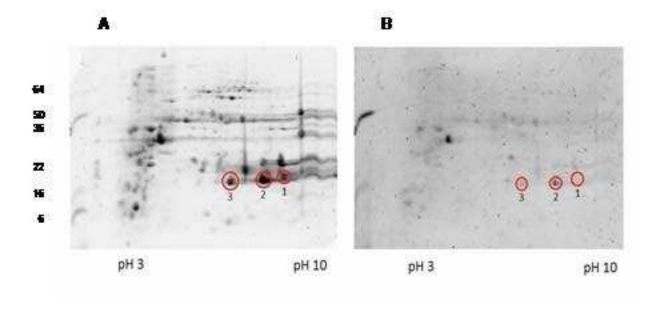


Figure 2

MASKATLLLAFTLLFATCIARHQQRQQQQQQQQQLQNIEALEPIEVIQAEAGVTEIWDAYD
QQFQCAWSILFDTGFNLVAFSCLPTSTPLFWPSSREGVILPGCRRTYEYSQEQQFSGEGG
RRGGGEGTFRTVIRKLENLKEGDVVAIPTGTAHWLHNDGNTELVVVFLDTQNHENQLDEN
QRRFFLAGNPQAQAQSQQQQQRQPRQQSPQRQRQRQRQGQGQNAGNIFNGFTPELIAQSF
NVDQETAQKLQGQNDQRGHIVNVGQDLQIVRPPQDRRSPRQQQEQATSPRQQQEQQQGRR
GGWSNGVEETICSMKFKVNIDNPSQADFVNPQAGSIANLNSFKFPILEHLRLSVERGELR
PNAIQSPHWTINAHNLLYVTEGALRVQIVDNQGNSVFDNELREGQVVVIPQNFAVIKRAN
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PSFSRGQGIRASR

Figure 3

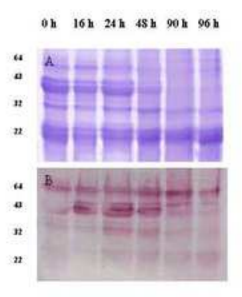


Table 1

3 Proteins identified in extracts from 16-h-imbibed sunflower seeds by peptide mass

4 fingerprinting.

Spot	Protein identification	Species	Accession number UniProt	Molecular mass Exper/Theor (kDa)	Peptides matched	Coverage (%)	Score
1	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	19.97/20.98 (β)	3	5.9	154
2	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	19.97/20.98 (β)	15	27.6	113
3	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	19.46/20.98 (β)	13	21.3	86

Table 2

Proteins identified in the eluted fraction of phosphoprotein affinity chromatography from seed extracts by peptide mass fingerprinting

Band	Protein identification	Species	Accession number UniProt	Molecular mass Exper/Theor (kDa)	Peptides matched	Coverage (%)	Score
1	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	29/20.98 (β)	10	18.5	34
2	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	25/20.98 (β)	5	9.1	110
3	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	24/20.98 (β)	18	26.4	290
4	11S globulin seed storage protein G3 precursor (α-β)	Helianthus annuus	P19084	22/20.98 (β)	8	15	202