Bioinformatic Analysis and Biophysical Characterization Reveal Structural Disorder in G0S2 Protein

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ABSTRACT: G0S2 is a small protein of 103 residues in length that is involved in multiple cellular processes. To date, several reports have shown that G0S2 functions by making direct protein–protein interactions with key proteins. In lipolysis, G0S2 specifically interacts with adipose triglyceride lipase, inhibiting its activity and resulting in lipolysis being downregulated. In a similar way, G0S2 also participates in the regulation of apoptosis, cell proliferation, and oxidative phosphorylation; however, information regarding G0S2 structural and biophysical properties is limited. In this work, we conducted a comparative structural analysis of human and mouse G0S2 proteins. Bioinformatics suggests the presence of a disordered C-terminal region in human G0S2. Experimental characterization by size-exclusion chromatography and dynamic light scattering showed that human and mouse G0S2 have different hydrodynamic properties. In comparison to the mouse G0S2, which behaves similar to a globular protein, the human G0S2 shows an elongated conformation, most likely by displaying a disordered C-terminal region. Further analysis of hydrodynamic properties under denaturing conditions suggests the presence of a structural element in the mouse protein that undergoes an order to disorder transition at low urea concentration. Structural analysis by circular dichroism revealed that in native conditions, both proteins are mainly unstructured, showing the presence of beta sheet structures. Further analysis of CD data suggests that both proteins belong to the premolten globule family of intrinsically disordered proteins. We suggest that the intrinsic disorder observed in the G0S2 protein may facilitate its interaction with multiple partners in the regulation of cellular metabolism.

INTRODUCTION

The G0/G1 switch gene 2 (G0S2) was first identified in blood mononuclear cells. The gene was differentially expressed during the drug-induced cell cycle transition from the G0 to G1 phase.1 The G0S2 gene encodes a small protein of 103 residues in length that has only been identified in vertebrates; orthologs in organisms including nematodes or flies have not been reported.2 G0S2 is conserved among species, human and mouse proteins share 77.7% of global identity with major differences located at the C-terminal region.3 The best-known function of G0S2 is in lipolysis. The protein inhibits the hydrolase activity of adipose triglyceride lipase (ATGL) by interacting with the patatin domain of the enzyme. Inhibition is observed even in the presence of CGI-58, the natural co-activator of ATGL.4 The participation of G0S2 in direct protein–protein interactions with key proteins has been described in multiple cellular processes. In mitochondria, G0S2 specifically interacts with Bcl-2, preventing its interaction with Bax. In this way, G0S2 inhibits the formation of the antiapoptotic heterodimeric complex Bcl-2/Bax, favoring apoptosis.5 In hematopoietic stem cells, G0S2 interacts with a domain rich in RGG repeats of nucleolin. By doing this, G0S2 retains nucleolin in the cytosol and arrests cell proliferation.5 Furthermore, the interaction of G0S2 with F0F1-ATP synthase in cardiomyocytes stimulates ATP production during hypoxia.6

Recent studies have shown that G0S2 displays a lysophosphatidic acid acyltransferase activity. As a result, G0S2 promotes triacylglycerol (TG) accumulation in the liver in addition to its function as an inhibitor of TG hydrolysis by interacting with ATGL. Moreover, the G0S2 gene is upregulated by the lipogenic transcription factor liver X receptor α.7 In addition, the G0S2 gene has been found to be hypermethylated in certain types of cancer, indicating a possible role in cancer development.4,8

To date, no experimental three-dimensional G0S2 structure has been published and predictions of its structure by
homology modeling fail because of the lack of adequate structural homologues. Moreover, the progress made in elucidating its structure—function has focused on identifying minimal domains of interaction with other proteins, which it is proposed to involve the hydrophobic domain (HD), though there are currently no reports of further structural studies. In this work, we describe the results of a bioinformatic analysis and biochemical and biophysical characterization of human and mouse G0S2 proteins. Our results show that both proteins behave as intrinsically disordered proteins (IDPs) of the premolten globule like (PMG).

**RESULTS AND DISCUSSION**

**In Silico Analysis Suggests the Disordered Character of the Human G0S2 C-Terminal Region.** In *silico* analysis of human and mouse G0S2 sequences included the prediction of disordered regions using the GeneSilico MetaDisorder server.7 The analysis of the sequences indicates similar but not identical disorder trends for both proteins. Human and mouse G0S2 proteins share the same HD, which is predicted to be the most structured region of the protein (residues 27–42, Figure 1A,B), whereas the N- and C-terminal regions tend toward disorder. It is remarkable that in comparison with the C-terminal region of human G0S2 may display properties of an intrinsic disordered region. To gain additional insights into the possible conformation of the C-terminal region of G0S2, we analyzed the mean net charge and the mean hydrophobicity properties of G0S2 sequences as these properties have been used to distinguish IDPs from ordered proteins.16 Figure 1D shows that human G0S2C is located in a well-defined position that corresponds to IDPs; however, mouse G0S2C is located in the corresponding region for ordered proteins. These results suggest the presence of a disordered C-terminal region in the human G0S2 protein.

**G0S2 Expression and Purification.** To experimentally evaluate the disordered nature of G0S2, we cloned the cDNAs encoding the full-length human and mouse G0S2 proteins into the pET28 expression vector. Initial attempts to express the N-terminal 6XHis-tagged proteins were unsuccessful, despite modifying expression conditions such as temperature incubation, expression time, IPTG concentration, and BL21 strains
(results not shown). Therefore, we decided to express the recombinant proteins as fusions with the SUMO protein from *S. cerevisiae* as previously reported. The SUMO-G0S2 proteins were successfully expressed and purified using nickel-nitrioltriacetic acid (Ni-NTA) affinity chromatography. The complete cleavage of the SUMO-G0S2 fusion proteins was achieved in reducing conditions (1–2 mM dithiothreitol (DTT)) with the HRV 3C protease. Soluble G0S2 proteins were further purified by ionic exchange and size-exclusion chromatography. The inset of Figure 2A shows the electrophoretic profile of both proteins after purification. The purity of both proteins, calculated with ImageJ software, was >95%.

**Figure 2.** Determination of Stokes radii for human and mouse G0S2 by gel filtration. (A) Analytical SEC chromatogram; the red and blue curves represent the elution profile of human and mouse G0S2, respectively. The inset shows a sodium dodecyl sulfate–polyacrylamide gel electrophoresis of the purified human and mouse G0S2 proteins used in SEC analysis. (B) Calibration curve prepared by plotting the *K*~AV~ values of standard proteins (black dots) versus their corresponding *R*~S~ value. The red and blue circles correspond to human and mouse G0S2, respectively.

**Hydrodynamic Analyses of Human and Mouse G0S2 Proteins.** Size exclusion chromatography (SEC) and dynamic light scattering (DLS) experiments were performed to obtain information on the hydrodynamic parameters related to the shape and size of both proteins. In analytical SEC experiments, human G0S2 eluted as a single peak (Figure 2A) with a retention volume corresponding to a Stokes radius (*R*~S~) of 18.8 Å. This value is 1.1 times larger than the theoretical value of 17.5 Å that can be calculated assuming that human G0S2 has a globular structure and a molecular weight of 11.3 kDa (Figure 2B). This difference resulted in being significant by means of a *t*-test of one sample (*P* = 0.0003). The experimental volume was 28.0 Å³, which was higher than the theoretical volume (22.4 Å³), while the experimental density of 0.4 kDa/Å³ was lower than the theoretical value (0.5 kDa/Å³), Table 1). The SEC elution profile for mouse G0S2 showed a single peak (Figure 2A) with an *R*~S~ of 17.5 Å having no significant difference (*P* = 0.0625) to the theoretical value of 17.4 Å, which was calculated with a molecular weight of 11.1 kDa (Figure 2B). The experimental values for volume (22.5 Å³) and density (0.5 kDa/Å³) were very similar to the theoretical values of 22.0 Å³ and 0.5 kDa/Å³, respectively (Table 1). These results suggest that the human G0S2 protein has a slightly extended structure compared to the more compact conformation observed in the mouse G0S2 protein. The extended conformation in human G0S2 could be explained in at least two possible ways. It may be due to a partially swelled tertiary structure, which would generate a globular conformation with larger dimensions, as in the case of a molten globule (MG). In this regard, the observed *R*~S~ value (18.8 Å) is smaller compared to the expected *R*~S~ value for an MG (19.9 Å). Alternatively, the extended conformation could also be attained by the presence of a disordered and extended region exposed to the solvent; in this case, the protein would have a globular domain with an extended segment that would generate an elongated conformation with the observed *R*~S~.

To gain further insights into the molecular shape of human and mouse G0S2 proteins, DLS measurements were subsequently carried out to calculate the hydrodynamic radius (*R*~h~) for each protein. The size distribution of protein samples showed the presence of a single species with an apparent *R*~h~ of 1.4 ± 0.03 and 1.7 ± 0.02 nm, human and mouse G0S2, respectively (Figure 3A). The hydrodynamic radius calculated for mouse G0S2 confirms that it has a compact and globular conformation as the *R*~h~ is in agreement with the *R*~S~ which is also 1.7 nm (Table 1). Interestingly, the *R*~h~ obtained for the human G0S2 protein was smaller than the calculated *R*~S~ value of 1.9 nm determined by SEC, indicating that this protein diffuses like a smaller protein in solution with an apparent size of 8 kDa. The apparent discrepancy could be explained as follows. DLS detects the translational diffusion motion in solution of macromolecules and estimates the corresponding translational diffusion coefficient (*D*), which is used to calculate the particle size using the Stokes–Einstein equation. The hydrodynamic dimensions are calculated by assuming that the diffusion particle is a sphere that diffuses at the same speed in all directions. Nonpherical particles (i.e., elongated proteins) can be represented by two axes of different lengths. These particles diffuse faster along one axis (the shorter) than the other (the longer) and as a result the size of the sphere calculated by DLS has a diameter that is shorter than the longer axis of the particle. Thus, an elongated protein will show an apparent *R*~h~ value that corresponds to a smaller size. In SEC, proteins are separated by their frictional coefficient (*f*) if the appropriate pore volume of the column has been selected. The calculated *R*~h~ is defined as the radius of a sphere that would have the actual *f* of the protein. The size of the sphere

"Calculated using the theoretical *R*~h~. *b*Calculated using the experimental *R*~h~."
determined by the $R_h$ is a good approximation to the size of a globular protein but elongated proteins usually show larger $R_h$ values. Thus, an elongated protein will show an apparent $R_h$ that corresponds to a larger size. Our SEC and DLS results suggest that the human G0S2 protein has an elongated shape with a globular domain of approximately 8 kDa. Similar results have been observed in the MG of the CaLB protein, whose radius calculated by DLS is smaller than that calculated by SEC. The elongated conformation of the human G0S2 could be explained by the presence of a disordered C-terminal region, as suggested by the bioinformatic analysis. Indeed, a theoretical molecular weight of 7.5 kDa can be calculated for human G0S2 by omitting the C-terminal region, which is in accordance with the observed $R_h$ value.

Given these $R_h$ differences in the native states of both proteins, we investigated their behavior under denaturing conditions; as both proteins are 103 residues in length and share an overall identity of 77.7%, we anticipated that unfolded proteins should have a similar $R_h$. Denaturation curves of human and mouse G0S2 were obtained in the presence of increasing urea concentrations, and at each point $R_h$ was estimated by DLS (Figure 3B). The denaturation curve for human G0S2 follows sigmoidal behavior with an increase in the experimental $R_h$; a single transition is observed from 2 to 6 M urea with midpoint transition $C_M$ of 4.4 M. The protein loses its structure and fully unfolds at 6 M urea, reaching an $R_h$ of 3.5 ± 0.06 nm; this value is in agreement with the expected $R_h$ value of 2.96 nm that can be calculated for an unfolded protein of 103 residues. The denaturation process of mouse G0S2 initiates with an $R_h$ of 1.7 ± 0.02 nm. A swelling process in which the side chains of the amino acids stop interacting with each other and start interacting with urea is observed from 0 to 2 M urea. Interestingly, when the protein is in 3 M urea its $R_h$ collapses to a value of 1.4 ± 0.01 nm. At higher concentrations, mouse G0S2 follows the same pattern observed for human G0S2 until it reaches a value of 3.4 ± 0.01 nm (Figure 3B). This result was interpreted as the loss of a structural element in the mouse protein, suggesting that at low urea concentration a structured region unfolds before the rest of the protein. The close similarity of mouse G0S2 $R_h$ value in 3 M urea (1.4 ± 0.01 nm) compared to the $R_h$ value observed for human G0S2 in native conditions (1.4 ± 0.03 nm) suggests that the labile structural element found in the mouse protein must be very similar in size and properties to that described for the human protein, therefore pointing to the C-terminal region of mouse G0S2 as the region that may contain the labile structural element.

Previous studies have shown that IDPs usually have regions that undergo disorder to order transitions that facilitate interaction with multiple partners. In order to identify a potential structural element in the C-terminal region of G0S2, we searched for molecular recognition regions (MoRFs) using the MoRFPred server. The prediction identified eight residues from 95 to 102 (ALSRLRQHA) in the mouse G0S2 sequence as a possible MoRF. Additionally, the Jpred4 server predicted an $\alpha$-helix structure for the same region (seven underlined residues). In the case of the human G0S2 protein, the region from residues 95 to 102 (ALSNRQHA) was also predicted as an MoRF but with a lower probability (six bolded residues out of eight) and only three residues were predicted to be in $\alpha$-helix conformation. Additionally, a hydrophobic cluster analysis (HCA) was performed. HCA provides a secondary helical representation of protein sequences in which hydrophobic clusters are plotted along the sequence. This approach has proven useful in identifying coiled coil regions, regions with biased composition, and regions with potential for induced folding via disorder–order transitions. HCA analyses of human and mouse G0S2 distinguish three segments that overlap to the N-terminal region, the HD and the C-terminal region. The N-terminal region displays short hydrophobic clusters; the main hydrophobic cluster is found within the HD, while the C-terminal region (resides 50 to 103) shows isolated hydrophobic residues with a couple of short hydrophobic clusters of two or three residues in length (Figure 4A,B). This is in agreement with the fact that long regions devoid of clusters correspond to disordered regions and small clusters within disordered regions correspond to putative MoRFs. The most striking difference is in residues 95 to 102 of mouse G0S2, which show “mosaic clusters” of leucine residues. Such clusters have been associated with secondary structure elements on the surface of proteins, thus suggesting the presence of a structural element in the C-terminal region of mouse G0S2 (Figure 4B). As a result of these analyses, the prediction of an MoRF region of eight residues, of which seven are predicted to be in an $\alpha$-helix conformation, as well as the presence of “mosaic clusters” of leucine residues, allow us to suggest that residues 95 to 102 of mouse G0S2 are part of the structural element that unfolds at low urea concentrations. In the case of human G0S2, the lower probability of MoRF, the prediction of a shorter $\alpha$-helix, and the absence of “mosaic clusters” support the disordered character of its C-terminal region.

Secondary Structure Content Analysis Reveals That G0S2 Is Largely Unstructured. To gain more insight into the structural content of G0S2, we used circular dichroism (CD) spectroscopy in the far UV region. The human and mouse G0S2 CD spectra exhibit properties previously observed for IDPs. They show a strong minimum signal close to 200 nm ($-11.67 \times 10^{2}$ and $-12.05 \times 10^{2}$ deg·cm$^{-1}$·dmol$^{-1}$, human and mouse G0S2, respectively) and a small negative contribution close to 222 nm ($-2.55 \times 10^{3}$ and
−1.48 × 10−3 deg cm² dmol−1, human and mouse G0S2, respectively) (Figure 5A). These characteristics have been

associated with the presence of residual structures in the order of 10−20%. The analysis of CD data using the BestSel algorithm3 indicates that human and mouse G0S2 proteins are mainly unstructured, showing content of ∼55% of coil, ∼25% of β-strands, ∼19% of β-turns, and virtually no contribution of α-helices (Table 2). The high content of coils confirms the tendency toward disorder suggested by the bioinformatic analysis (Figure 1B). The content of β-structures is in agreement with the secondary structure prediction of G0S2. Previous reports have suggested the presence of two alpha-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a hydrophobic sequence with the potential to assume β-sheet conformation. A detailed examination using the JPred4 and the fast estimator of latent potential to assume α-helices separated by a

Figure 5. Far-UV CD spectra of human and mouse G0S2. (A) CD spectra of human (red) and mouse (blue) G0S2 were recorded at 22 μM protein concentration. (B) Double-wavelength plot showing ellipticity [θ] at 200 and 222 nm of a set of coil-like (white circles) and PMG-like (gray circles) proteins. The positions of human and mouse G0S2 are indicated by red and blue circles, respectively.

Table 2. Summary of Human and Mouse G0S2 Secondary Structure Content As Estimated by the BeStSel Algorithm

| type of structure | hG0S2 | mG0S2 |
|------------------|-------|-------|
| Helix            | 0.4   | 0     |
| Antiparallel     | 24.8  | 25.8  |
| Parallel         | 0     | 0     |
| Turn             | 19.1  | 19.6  |
| Other (coil)     | 55.7  | 54.6  |
| Total            | 100   | 100   |

MATERIALS AND METHODS

In Silico Analysis. The following servers were used online with the default parameters. Protein disorder prediction was carried out using the GeneSilico MetaDisorder server9 (http://genesilico.pl/metadisorder/). Composition Profiler10 (http://www.cprofiler.org) was used to analyze the amino acid distribution. Charge-hydrophathy plot analyses16 were generated using PONDR (http://www.pondr.com). HCA28 was performed at (http://mobyle.rpbs.univ-paris-diderot.fr/cgi-bin/portal.py?form=HCA#forms::HCA). Protein secondary structure prediction was done by Jpred4 (https://www.compbio.dundee.ac.uk/jpred/) and FELLs (http://old.protein.bio.unipd.it/fells/) servers.29

Protein Expression and Purification. The coding DNA for human (NP_056529.1) and mouse (NP_032085.1) G0S2 and S. cerevisiae SUMO protein residues 1−97 (6xHis-SUMO-HRV3C) was acquired optimized for bacterial expression from Integrated DNA Technologies. The SUMO fragment was cloned into a pET28a expression vector, followed by the insertion of the G0S2 fragment to obtain the pET28SUMO-hG0S2 and pET28SUMO-mG0S2 constructs. Escherichia coli BL21 (DE3) competent cells were transformed with the recombinant vector and cultured for 4 h at 37 °C. Protein expression was induced with 1 mM IPTG at 28 °C overnight. The cells were harvested and sonicated at 4 °C in lysis buffer [50 mM Tris (pH 8.0), 300 mM NaCl, and 10 mM imidazole]. The supernatant was loaded onto a Ni-NTA column equilibrated with the lysis buffer. Fusion proteins were eluted
in one step with the lysis buffer supplemented with 500 mM imidazole. 6xHis-SUMO fusion was removed with HRV 3C protease overnight at 4 °C. Human G0S2 was then loaded into a HiTrap-Q HP column (GE Healthcare) in buffer A [50 mM Tris buffer (pH 8.0), 0.5 mM ethylenediaminetetraacetate (EDTA), and 1 mM DTT], and mouse G0S2 was loaded into a HiTrap-S HP column in buffer B [50 mM HEPES (pH 8.2), 0.5 mM EDTA, and 1 mM DTT]. Eluted proteins were concentrated (Amicon Ultra-15, Merck) and further purified through a Superdex-75 10/300 GL column (GE Healthcare) with buffer C [20 mM Tris (pH 8.0), 150 mM NaCl, 0.5 mM EDTA, and 1 mM DTT]. Protein concentration was estimated at 205 nm; extinction coefficient values (ε205nm) of 29.70 and 30.23 were estimated directly40 for human and mouse G0S2, respectively.

**Analytical Size-Exclusion Chromatography.** Protein samples of 100 μL at 1 mg/mL were injected independently on a Superdex 75 10/300 GL column (GE Healthcare), equilibrated with SEC buffer [20 mM Tris–HCl pH 8.0 and 150 mM NaCl] at a flow rate of 0.5 mL/min. The column was calibrated with protein markers: ovalbumin (44 kDa), myoglobin (17 kDa), insulin (5.8 kDa), and vitamin B12 (1.35 kDa). Stokes radii (Rg) were calculated by means of eq 1.20

\[
\log(R_g) = -(0.204 \pm 0.023) + (0.357 \pm 0.005) \log(MW)
\]

The total volume of the column (Vt) was 24 mL and the void volume (V0) of 8.2 mL was calculated using thyroglobulin. The gel phase distribution coefficients (KAV) for the standard proteins were determined using eq 2,41 where Vt is the elution volume of the protein.

\[
K_{AV} = \frac{V_k - V_0}{V_t - V_0}
\]

**Dynamic Light Scattering.** DLS experiments were performed at 25 °C using an APS20000 system (Malvern Instruments). All hydrodynamic radii (Rg) measurements were made at 1 mg/mL protein concentration in buffer D [20 mM Tris (pH8.0) and 1 mM beta-mercaptoethanol]. For denaturation curves, proteins were incubated with increasing concentrations of urea overnight at room temperature prior to DLS measurements being taken. All values are the mean of three independent measurements.

**CD Spectroscopy.** CD measurements were made on a JASCO J-815 spectropolarimeter (Jasco Inc., Easton, MD) equipped with a PDR-425S Peltier temperature controller cell. CD spectra were recorded from 194 to 250 nm using a 0.1 cm path length cell. Protein samples were prepared in buffer E [20 mM potassium phosphate (pH 8.0) and 20 mM NaCl] at 0.25 mg/mL. Ellipticity is reported as mean ellipticity per residue \( [\theta] \). Secondary structure content was estimated from the CD data (range 200–250 nm) using BestSel (http://bestsel.eltc.hu/).33

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**Notes**

The authors declare no competing financial interest.

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**NOMENCLATURE**

G0S2, G0/G1 switch gene 2; IDP, intrinsic disorder protein; PMG, premolten globule; SEC, size-exclusion chromatography; DLS, dynamic light scattering; CD, circular dichroism; MoRFs, molecular recognition regions; HCA, hydrophobic cluster analysis

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