The Actin Binding Protein, Fesselin, is a Member of the Synaptopodin Family ,#,†

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Abstract

Fesselin is a natively unfolded protein that is abundant in avian smooth muscle. Like many natively unfolded proteins, fesselin has multiple binding partners including actin, myosin, calmodulin and α-actinin. Fesselin accelerates actin polymerization and bundles actin. These and other observations suggest that fesselin is a component of the cytoskeleton. We have now cloned fesselin and have determined the cDNA derived amino acid sequence. We verified parts of the sequence by Edman analysis and by mass spectroscopy. Our results confirmed fesselin is homologous to human synaptopodin 2 and belongs to the synaptopodin family of proteins.

Keywords

Fesselin; synaptopodin; myopodin; actin binding protein; actin polymerization; Ca2+-calmodulin regulation

Fesselin is an actin binding protein [1] that is rich in avian smooth muscle. Fesselin binds to both F-actin and to G-actin. When added to G-actin, fesselin induces rapid polymerization by increasing the rate of nucleation [2]. The rate of actin polymerization is inhibited by Ca2+-calmodulin to a rate that is lower than that of actin alone [3]. When added to F-actin fesselin initiates the formation of actin aggregates that sediment at low speed [1]. Fesselin inhibits actin activation of myosin S1 ATPase activity [4]. That inhibition is not reversed by Ca2+-calmodulin. Fesselin is natively unfolded [5] with little secondary or tertiary structure when in pure form at neutral pH and physiological ionic strength. Like other natively unfolded proteins fesselin has several binding partners and gains secondary structure when bound to one of those partners (calmodulin). In addition to binding actin and calmodulin [3], fesselin also binds to myosin [4] and α-actinin [6]. These abilities of fesselin to polymerize actin, organize actin

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1The turkey fesselin sequence reported in this paper has been deposited in GenBank under the name Turkey Synaptopodin 2 (GenBank accession nos. EU086520, ABU55374).

2Myopodin was first described as a unique gene product (Weins et al. 2001) and was later shown to be an alternative splicing product of the synaptopodin 2 gene.

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filaments and bind to other actin associated proteins suggest a role of fesselin in organizing cellular actin. This hypothesis is supported by evidence that fesselin is localized in smooth muscle dense bodies [7].

Because actin binding proteins tend to be highly conserved it is likely that proteins similar to fesselin are found in mammals [8]. In an earlier analysis of 7 fesselin peptides (about 8% of the total fesselin sequence) we observed that three fragments were unique and 4 others had 57–70% identity to synaptopodin [1]. Synaptopodin is the eponym of a family of actin binding proteins that is formed by 3 protein subgroups: synaptopodin, the synaptopodin 2 proteins, to which myopodin belongs, and the synaptopodin-2 like proteins. Both fesselin and synaptopodin are rich in proline and have high isoelectric points [1,9]. Synaptopodin family members also bind to actin [9,10] and α-actinin [11,12]. Synaptopodin has been linked with the organization of the cytoskeleton [13].

Synaptopodin family members are important since they have been associated with defects of the spine apparatus and learning deficiencies in knock out mice [14]. These proteins are also linked to some cancers. Myopodin suppresses cancers of bladder and prostate [15,16] and its absence has been used as an indication for cancer staging.

Establishing a link between fesselin and synaptopodin family members is important because the ease in purification of avian fesselin facilitates biochemical and biophysical analyses of this protein. As such, fesselin could be a useful paradigm for the synaptopodin family. We have now cloned fesselin. From the derived sequence and from direct sequencing of the protein we now report that avian fesselin is homologous to mammalian synaptopodin 2.

Materials and Methods

Protein-Sequencing

Sequencing was performed by Edman degradation [17]. Proteins were separated on SDS-gels and transferred on Sequi-Blot™ PVDF Membrane (BIO-RAD) in a tank blot apparatus [18]. To obtain specific peptide fragments, the protein was digested with trypsin directly on the PVDF membrane [19]. The peptides obtained were separated by high pressure liquid chromatography (HPLC), on a Hypersil C18 BDS 3-m LC-Packings column (150.1.0 mm; BIA, Bensheim, Germany) and a 130 A HPLC separation system (Applied Biosystems, Weiterstadt, Germany) with 0.1% trifluoric acid (TFA) as solvent A and 80% acetonitrile in 0.085% TFA as solvent B. The HPLC-separated fragments were sequenced on Polybrene-treated filters, using a Procise 494 A protein sequencer from Applied Biosystems. MALDI-TOF fingerprint analysis was performed by Dr. Monica Linder (Giessen, Germany). Proteins were precipitated with 8 volumes acetone at −20°C and then washed 3 times with acetone (−20°C) prior to shipping. Peptides were identified using the program Mascot Search Results from MATRIX SCIENCE Inc., Boston, MA, and the program ProteinProspector v4.0.8, http://prospector.ucsf.edu/.

RNA-isolation, RT-PCR, cloning, and DNA-sequencing of turkey fesselin

Total-RNA was isolated from turkey gizzard tissue using an RNeasy Kit (Qiagen, Hilden, Germany). The RT-reaction was performed using dT_{18}-oligos and Superscript II or Superscript III (Invitrogen, Carlsbad, CA). For the PCR reactions oligo nucleotides were selected based on the nucleotide sequence for a predicted protein similar to synaptopodin 2 from gallus gallus (GenBank accession no. XM_426324). Oligo nucleotides used were as following: 5’ primer 5’ AGA GAG AAT TCT CCT TCT CTG CCC TCC TTT G 3’ (bp 1316–1336), 3’ primer 5’ AGA GAG GAT CCT CAC CTG GAA TCC ATG GAC ATT ATA TCA G 3’ (bp 4099–4073). The 5’ and 3’ ends of the mRNA were obtained using a 5’/3’ RACE Kit, 2nd...
Protein preparations

Turkey gizzard fesselin was purified as described by Leinweber et al. [1].

Western blot

Snap frozen tissues were ground under liquid nitrogen to a fine powder and suspended in a buffer containing 8 M urea, 0.1 M Tris pH 6.8, 2% SDS, 0.035M dithiothreitol, and protease inhibitor cocktail (SIGMA P2714) equivalent to 0.1 mg USP pancreatin/ml. The suspended tissue was incubated for 1 hour on ice, and clarified by centrifugation. A Bradford protein assay was used to determine the protein concentration in the supernatants. 50 µg proteins per lane were electrophoresed on a 10% SDS gel, transferred on nitrocellulose membrane, and probed with anti fesselin antibodies.

Results and Discussion

In our first description of turkey gizzard fesselin [1] we sequenced 7 short peptides of that new protein. We noted that 4 of these peptides had identities ranging from 57 to 70% with the previously published synaptopodin sequence [9]. Those regions of homology represented approximately 4% of the total fesselin sequence. While our limited sequence information showed some similarity to the synaptopodin family the distribution of fesselin is different from that reported for any of the synaptopodin family member proteins. Fesselin is abundant in avian smooth muscle tissue. Synaptopodin is found in kidney and brain [9]. Synaptopodin mRNA, but not the gene product, has been found in pancreas [20]. The splicing product of synaptopodin 2, myopodin, is expressed in many tissues and is particularly rich in skeletal muscles with smaller amounts in smooth muscles [10,16]. None of the synaptopodin family members is abundant in mammalian smooth muscle.

To explore the possibility that fesselin is a member of the synaptopodin family we have determined the sequence of fesselin both at the protein and cDNA levels. Knowing this relationship is important for determining the function of fesselin and synaptopodin. More information is available on the biochemistry of the more readily purified fesselin while much is known about the cell biology of synaptopodin family members.

Avian fesselin is isolated as two polypeptide chains (103 and 79 kDa) having similar properties [1]. We analyzed the sequence of purified 103 kDa natural fesselin form by both Edman and MALDI-TOF analyses to explore the relationship to the synaptopodin family. Results from Edman analysis of internal peptides are shown in Table 2. Of the 116 residues analyzed, 112 were identical to predicted avian synaptopodin 2 (96.5% identity); 3 residues were conservative substitutions. The regions of homology extended from amino acid residues 498 to 1092 of the gallus gallus sequence (GenBank accession no XM_426324). Figure 1 shows the alignment of the cDNA derived amino acid sequence from turkey fesselin (GenBank accession nos. EU086520, ABU55374) with predicted synaptopodin 2 from gallus gallus (GenBank accession no. XM_426324). Amino acid residues that were confirmed by Edman analysis are shown as bold letters in that figure. Note that we were unable to identify the N-terminal residue by Edman analysis.
We used MALDI-TOF MS analysis to compare 66 tryptic peptides of the 103 kDa fesselin polypeptide with the predicted synaptopodin 2 like protein from *gallus gallus* (GenBank accession no. XM_426324). The results of the MALDI-TOF analyses are underlined in Figure 1. The 66 peptides were distributed over the whole *gallus gallus* sequence allowing us to confirm 66% of the cDNA-derived sequence. The carboxyl terminus identified by mass spectroscopy from the 103 kDa polypeptide corresponded to the C-terminus of the predicted *gallus gallus* sequence. However, the first N-terminal residue identified by these peptides was equivalent to amino acid number 268 of the predicted *gallus gallus* sequence.

We confirmed the sequence of the 103 kDa fesselin by an analysis of the cDNA sequence. The region of known amino acid sequence was amplified by RT-PCR. Separation of the RT-PCR products performed on turkey gizzard RNA resulted in a band at about 2.8 kb (not shown) that, when sequenced, was found to be equivalent to nucleotides 941–3718 of predicted synaptopodin 2 from *gallus gallus*.

The N- and C-terminal regions of fesselin were defined using 5’ and 3’ RACE as described in Materials and Methods. We observed different N-termini with 3 independent 5’ RACE studies. Our earliest detected upstream nucleotides corresponded to amino acid residues 174 or 263 of the chicken sequence. This means that the likely start sites in mature 103 kDa fesselin are either methionine 194 or methionine 268. With MALDI-TOF analysis our first detected peptide corresponded to amino acid residues Met268–Arg289 indicating this is the most likely start point of mature fesselin. The molecular mass predicted from the start site at methionine 268 (107.702 kDa) is in agreement with our earlier estimate (103 kDa) [1].

The presumed synaptopodin 2 of the domestic chicken does not have a methionine residue at position 268. In fact, residue 268 comes in them middle of an exon of the chicken. Yet the apparent mobility of turkey and chicken fesselin’s on SDS gels are the same (see Figure 2). We must consider the possibility that the 103 kDa fesselin band is the product of proteolytic processing. In some preparations of fesselin we have detected larger forms of fesselin. Figure 2 shows extracts of both turkey and chicken gizzard muscle probed by anti-fesselin antibodies. The two higher forms have approximate apparent molecular weights of 140 and 160 kDa. We confirmed that these higher bands are in fact derived from fesselin by MALDI-TOF fingerprint analysis. The most N-terminal sequence identified to this point is residues 256 to 264. Further analyses of the function of these larger fesselin forms may lead to as yet unidentified binding partners. These larger fesselin forms may also provide clues to processing that gives rise to the major forms of fesselin that are isolated.

The major fesselin forms having molecular masses of 79 and 103 kDa lack a PDZ domain that is present in the predicted synaptopodin 2 sequence from chicken (aa 7–85). PDZ domains are small globular protein-protein interacting modules consisting of six β-strands and two α-helices. The majority of the interactions consist of the binding to a carboxyl terminal pentapeptide of the target protein or the binding of C-terminus mimicking motifs. The structure of these highly specialized domains does not change significantly upon protein binding [21]. The larger fesselin precursors may therefore have interactions with proteins that are to this point unknown.

In summary, the 103 kDa fesselin band is an avian gene product of predicted synaptopodin 2 (GenBank accession no. XM_426324). We have shown this by comparing results obtained from analyses of the isolated protein (Edman and MALDI-TOF MS) and of the primary sequence derived from the cDNA of fesselin with the sequence of predicted avian synaptopodin 2 [XM_426324]. Further evidence of identity between fesselin and synaptopodin 2 is that the first 21 hits in a BLAST P search [22] performed with the cDNA derived amino acid sequence of turkey fesselin were synaptopodin 2 protein sequences with an E value of 0 indicating
identity. Turkey fesselin has 60% identity and 72% similarity with human synaptopodin 2 (GenBank accession no. NP_597734). Fesselin is also 60% identical to human myopodin (GenBank accession no. CAB51856), an alternative splicing product of synaptopodin 2. In comparison, the similarity of fesselin to human synaptopodin (GenBank accession no. AAQ07402) is approximately 43%. This shows fesselin is homologous to mammalian synaptopodin 2. Furthermore, like synaptopodin 2, fesselin is rich in proline and contains multiple PXXP motifs. These PXXP motifs could function as potential peptide ligand motifs for SH3 domain containing proteins. Fesselin, like synaptopodin 2, contains one PPXY motif. These motifs function often as ligand peptides for proteins with WW domains as found in dystrophin [23].

Acknowledgements

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References

7. Renegar RH, Chalovich JM, Leinweber BD, Zary JT, Schroeter MM. Localization of the actin-binding protein fesselin in chicken smooth muscle. submitted


Abbreviations

EDTA, ethylenediaminetetraacetic acid; EGTA, ethylene glycol-bis(β-aminoethyl ether)-N,N,N′,N′-tetraacetic acid; MOPS, 3-(N-Morpholino)-propanesulfonic acid.
Figure 1.
Alignment of the cDNA derived amino acid sequence of 103 kDa turkey fesselin (GenBank accession nos. EU086520, ABU55374) with that of predicted synaptopodin 2 from \textit{gallus gallus} (GenBank accession no. XM_426324) starting with residue 268 in the chicken sequence. MALDI TOF fingerprint results are underlined. Identical residues are indicated by : and similarities are shown by +. Edman analyses are in bold letters.
Figure 2.
Western Blot of gizzard homogenate from turkey (lane a) and chicken (lane b) probed with polyclonal affinity purified anti-fesselin antibodies raised against gel excised fesselin purified from turkey.
<table>
<thead>
<tr>
<th>Oligo Nucleotide</th>
<th>Sequence</th>
</tr>
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<tbody>
<tr>
<td>5' RACE</td>
<td></td>
</tr>
<tr>
<td>1st primer</td>
<td>5’ ACC TGG AAT CCA TGG ACA TTA TAT CAG 3’</td>
</tr>
<tr>
<td>2nd primer</td>
<td>5’ TCA CCT TCT TCA CCC TCG TC 3’</td>
</tr>
<tr>
<td>3rd primer</td>
<td>5’ AAA CAT GAG GGC CCC TTT AC 3’</td>
</tr>
<tr>
<td>4th primer</td>
<td>5’ GTT GCC CTG TGC TTG AGA G 3’</td>
</tr>
<tr>
<td>3' RACE</td>
<td></td>
</tr>
<tr>
<td>1st primer</td>
<td>5’ GAA AGC AAC CGG GTG TTA CAG 3’</td>
</tr>
</tbody>
</table>
### Table 2

Peptides of turkey fesselin determined by Edman degradation \( ^a \)

<table>
<thead>
<tr>
<th>Protease</th>
<th>Peptide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trypsin</td>
<td>TGILQEAK (681–688); STSKPMFSFK (691–700); APSF-APASPQAAYP (789–803); GAQLFAK (868–874); FARRHSRMEK (872–892); YVVDSDTVQAN (882–892); CSLSPLRPIPI (1082–1092)</td>
</tr>
<tr>
<td>Glu C</td>
<td>ACNFMOS (743–749); TVQANMARASSP (888–899); SNVRAPPVVA (911–920)</td>
</tr>
<tr>
<td>Arginase C</td>
<td>RORMDOITAEQEE (498–510); EAPKVPSPNAPSL (701–714)</td>
</tr>
</tbody>
</table>

\( ^a \)The table does not include peptides that were published earlier [1]

\( ^b \)The numbers correspond to the position in the predicted synaptopodin 2 from *gallus gallus* (GenBank accession no. [XP_426324](https://www.ncbi.nlm.nih.gov/protein/XP_426324)), underlined residues are tentative.