The Olfactomedin Domain from Gliomedin Is a β-Propeller with Unique Structural Properties*

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Background: Gliomedin contains an olfactomedin (OLF) domain, for which structural information has been lacking.

Results: The crystal structure of the OLF domain from gliomedin was solved.

Conclusion: The OLF domain is a five-bladed β-propeller with broken symmetry.

Significance: The results provide a structural basis for gliomedin function and shed light on mutations in the human OLF family.

All members of the olfactomedin (OLF) family have a conserved extracellular OLF domain, for which a structure has not been available. We present here the crystal structure of the OLF domain from gliomedin. Gliomedin is a protein expressed by Schwann cells in peripheral nerves, important for the formation of the nodes of Ranvier. Gliomedin interacts with neuronal cell adhesion molecules, such as neurofascin, but the structural details of the interaction are not known. The structure of the OLF domain presents a five-bladed β-propeller fold with unusual geometric properties. The symmetry of the structure is not 5-fold, but rather reveals a twisted arrangement. The conserved top face of the gliomedin OLF domain is likely to be important for binding to neuronal ligands. Our results provide a structural basis for the functions of gliomedin in Schwann cells, enable the understanding of the role of the gliomedin OLF domain in autoimmune neuropathies, and unravel the locations of human disease-causing mutations in other OLF family members, including myocilin.

Gliomedin is a protein expressed by myelinating Schwann cells that is important for the formation of the nodes of Ranvier and clustering of Na⁺ channels on the axonal plasma membrane at the node (4–6). Gliomedin contains an extracellular olfactomedin (OLF) domain, which is able to induce channel clustering (7, 8). Gliomedin molecular ligands include neurofascin and NrCAM (neuronal cell adhesion molecule) (5, 9), and loss of neuronal neurofascin also results in depletion of gliomedin from the nodes of Ranvier (10). Gliomedin is a type II transmembrane protein, and in addition to its C-terminal OLF domain, it harbors an extracellular collagen-like domain, which mediates gliomedin trimerization (8). Gliomedin homologs have also been identified in invertebrates (11). In vivo, gliomedin is processed proteolytically, and the extracellular OLF domain can be released either with or without the collagen domain (7, 8). The OLF domain is also glycosylated (8), although the exact sites and nature of glycosylation have not been determined.

The ~250-residue OLF domain was first discovered in the early 1990s (12). All members of the OLF protein family contain an extracellular OLF domain (13). OLF domains are mediators of extracellular protein interactions, and they play roles in diverse processes, such as nervous system development, intercellular adhesion, and cell cycle regulation (14). Although the structure of the OLF domain has remained unknown, it can be predicted to consist largely of β-structure. The OLF domain of myocilin was suggested to form a six-bladed β-propeller structure (15), but no experimental data have backed up this prediction.

Due to its direct disease linkage, the most widely studied OLF family member is myocilin, mutations in which cause primary open-angle glaucoma in humans (16–18). Myocilin is also involved in myelination, especially in the optic nerve (19–22). The sequence identity between the OLF domains of gliomedin and myocilin is only 22%, however, suggesting that they do not carry out overlapping functions. An interaction between gliomedin and myocilin has also been reported (20).

We determined the high-resolution crystal structure of the OLF domain from gliomedin. The structure presents an unusual β-propeller fold, and it can be used both to understand
the involvement of gliomedin in neurological development and disease and to obtain information on disease-related mutations in related OLF family members, such as myocilin.

**EXPERIMENTAL PROCEDURES**

**Protein Expression and Purification**—The OLF domain of rat gliomedin (residues 260–543; sequence numbering includes the signal sequence) was recombinantly expressed in baculovirus-infected insect cells (23). Purification consisted of nickel-nitrilotriacetic acid affinity chromatography, His tag cleavage using tobacco etch virus protease, and size-exclusion chromatography (23). The OLF domain was predicted by sequence analysis to lie at the very C terminus, whereas the N-terminal boundary of the domain was difficult to pinpoint. The domain selected for expression and structural studies ranged from the end of the collagen-like domain close to the C terminus. This construct provided a good yield of crystallizable recombinant protein.

**Biophysical Characterization**—CD spectroscopy was used to confirm the folding state of the purified gliomedin OLF domain. Spectra were measured on an Applied Photophysics Chirascan-plus instrument at +20 °C in a buffer containing 1.8 mM Tris (pH 7.5) and 7 mM NaCl. The protein concentration was 0.5 mg/ml, and the cuvette path length was 0.5 mm.

Synchrotron small-angle x-ray scattering (SAXS) data were obtained on the SWING beamline at the SOLEIL Synchrotron (Paris, France). The gliomedin OLF domain (6 mg/ml) was injected onto an Agilent SEC-3 size-exclusion column using an Agilent HPLC system and eluted with 0.1 M Tris (pH 7.5) and 400 mM NaCl. The eluate was directed to the SAXS flowthrough capillary cell, and scattering data were collected online. Frames corresponding to the main peak were selected and averaged using FoxTrot. Following data processing, further analyses were performed with the ATSAS package (24). GASBOR (25) was used for ab initio model building. CORAL (24) was used in conjunction with the crystal structure to build the fragments not visible in the crystal.

**Crystalization and Data Collection**—Highly concentrated pure gliomedin OLF domain (23 mg/ml) in 50 mM Tris (pH 7.5) and 100 mM NaCl was used for crystallization. After 3 days of incubation at 293 K, crystals were observed in 0.1 M CHES (pH 7.5) and 7 mM NaCl. The protein concentration was 0.5 mg/ml, and the cuvette path length was 0.5 mm.

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Crystal Structure of the Olfactomedin Domain—The structure was solved using the single-wavelength anomalous dispersion protocol of Auto-Rickshaw, the European Molecular Biology Laboratory (EMBL) Hamburg automated crystal structure determination platform (27). The input diffraction data were prepared and converted for use in Auto-Rickshaw using programs of the CCP4 suite (28).

Estimated substructure structure factor amplitude values were calculated using SHELXC (29). Based on an initial analysis of the data, the maximum resolution for substructure determination and initial phase calculation was set to 2.61 Å. Heavy atoms were found with SHELXD (30), and the correct hand for the substructure was determined using ABS (31) and SHELXE (32). Initial phases were calculated after density modification using SHELXE (32). The 2-fold non-crystallographic symmetry operator was found using RESOLVE (33). Density modification, phase extension, and non-crystallographic symmetry averaging were performed with dm (34). Two chains of 242 residues each were thereafter automatically built using ARP/wARP (35, 36).

The automatic pipeline further refined this partial model to 2.0 Å resolution with the platinum derivative data using CNS (37), Refmac5 (38), and PHENIX (39). The partially refined structure was then used as a template for molecular replacement against a native high-resolution data set in a different space group. The structure was further refined using PHENIX and manually built using Coot (40). The final coordinates and original structure factors were deposited in the Protein Data Bank with codes 4D77 (orthorhombic) and 4D7C (monoclinic).

**Homology Searches and Structure Analysis**—Homologous structures were searched using PDBbind (41) and SALAM (42). For structure analysis and visualization, the programs PyMOL, UCSF Chimera (43), WEBnrg (44), CCP4mg (45), and ConSurf (46) were used. A homology model for the myocilin OLF domain was constructed based on sequence alignment between myocilin and gliomedin using SWISS-MODEL (47).

**RESULTS**

**Crystal Structure of the Gliomedin OLF Domain**—Gliomedin contains an N-terminal transmembrane domain, a collagen-like segment, and a C-terminal OLF domain (Fig. 1A). The structure of the rat gliomedin OLF domain was solved by single-wavelength anomalous dispersion phasing using a crystal derivatized with platinum, and the structure was refined to 1.49 Å resolution using native data (Table 1). The OLF domain structure is composed of a five-bladed β-propeller (Fig. 1B), and each blade contains four β-strands (strands a–d). The overall dimensions of the OLF domain are ≈30 × 40 × 40 Å. Like other β-propellers, the OLF domain is disc-shaped. The propeller circle is closed by insertion of the first N-terminal β-strand into the framework of the last propeller blade (Fig. 1B), which is a common mechanism in β-propeller proteins, also termed Velcro closure (48). The structure also contains three short α-helices. The longest of these helices is located in a loop between β-strands c and d of blade 1. Interestingly, normal mode analysis of the structure suggests that this helix is highly mobile, and there may be significant motions between blades 1 and 5 (data not shown). In general terms, normal mode analysis allows the identification of potential low-frequency, large-amplitude,
concerted motions in a protein structure. A schematic representation of the OLF domain topology is shown in Fig. 1C.

The common fold of a β-propeller structure contains 4–10 antiparallel four-stranded β-sheets (49). Upon close inspection, the OLF domain five-bladed fold lacks a regular 5-fold symmetry and appears twisted. Indeed, blades 1–4 are actually arranged according to a hexagonal setting (Fig. 1C). The remaining blade 5, which contains both the N and C termini, actually takes up the space that would be filled by two blades in a six-bladed β-propeller. This arrangement is certainly unexpected and unusual, and it distinguishes the OLF domain from both five- and six-bladed β-propellers, also explaining difficulties in correctly predicting the OLF domain structure.

Structural homology searches using different algorithms with the gliomedin OLF domain produced rather poor hits to β-propeller structures, and both five- and six-bladed structures were detected. However, when only blades 1–4 were used in searches, better fits were obtained toward six-bladed structures. This further highlights the loss of 5-fold symmetry in the OLF domain and confirms the 6-fold symmetry obeyed by the first four blades. Superpositions on symmetric five- and six-bladed structures further proved this arrangement (Fig. 2).

Surface Properties—To obtain an insight into possible functionally important sites on the gliomedin OLF domain surface, we analyzed its surface properties. The electrostatic surface of the gliomedin OLF domain shows a high positive charge on the...
top face, whereas the bottom is more neutral or even negatively charged (Fig. 3A). The positive charge may be important in attracting binding partners of gliomedin.

The distribution of aromatic residues on the surface of the OLF domain (Fig. 3B) indicates a possible surface for protein interaction with several aromatic rings. For example, carbohydrate ligands would be candidates for interaction with the OLF domain.

The OLF domain structure was also refined in a monoclinic crystal form, and an apparent concerted movement of large side chains at the top face between the two crystal forms was observed (Fig. 3C). This indicates some flexibility in the structure, which could be important for interactions. A plasticity of the aromatic surface could allow binding of diverse protein and/or carbohydrate ligands.

Gliomedin is known to be glycosylated on its OLF domain (8), and a total of four possible N-glycosylation sites can be predicted to lie within the domain. The crystal structure indicates that all these possible sites of glycosylation lie on the surface. Remarkably, all the sites are on the “side” of the OLF domain, keeping the top face accessible (Fig. 3B).

**Solvent Cavity and Cation-binding Site**—In general, β-propellers have a central cavity, where, for example, enzymatic active sites can reside. The cavity or channel size increases as the number of blades gets higher (49). Gliomedin has a much smaller central cavity due to the disrupted symmetry. Furthermore, the top of this cavity is closed by a long loop (connecting strands 2d–3a), and it contains a number of water molecules and a bound cation (Fig. 3D).
A positively charged ion is present inside the gliomedin OLF structure, on the side of the water-filled cavity. The ion is well defined in electron density and coordinated by five oxygen atoms at distances of $\approx 2.30$ Å (Fig. 3E). Also taking into account the buffer composition in crystallization, this ion could be confidently built as Na$^+$. The ion was also validated by the WASP (50) and CheckMyMetal (51) servers, and clearly the best scores were obtained for the presence of Na$^+$. In particular, the Na$^+$-specific valence value of 1.1 calculated by WASP clearly indicates bound Na$^+$. Myocilin was shown to bind and be stabilized by calcium (52), and the D380A mutation abolished calcium binding, as well as the calcium-induced increase in protein stability. This mutation is also a disease-causing variant in myocilin (see below for more discussion on myocilin mutations).

Conservation in Gliomedin from Different Species—To get a better idea of putative functionally important parts of the OLF domain structure, sequences of gliomedin from different species were compared with regard to sequence conservation. In general, the least conserved regions include the outermost $\beta$-strands in the blades. On the other hand, in addition to the protein core, residues on the top surface are highly conserved and hence most likely involved in gliomedin function in vivo. This conservation coincides with the positive electrostatic potential and the location of the clusters of aromatic residues on the surface. The bottom face of the OLF domain is poorly conserved.

In addition to gliomedins, conservation was further checked for different representative OLF domain sequences (Fig. 4, B and C). Based on this analysis, 20 fully conserved residues were identified on OLF domains, of which three are glycine and seven are aromatic. It is obvious that these conserved residues are important for the unique structural aspects of the OLF domain fold due to their localization inside the fold. The top face of the OLF domain has none of these conserved sites, which is an indication of different binding preferences of OLF family proteins for their respective ligands.

**Figure 4.** Sequence conservation of OLF domains at the structural level. A, conservation in gliomedins from different species. Blue indicates high and yellow indicates poor conservation. The top surface (upper) is highly conserved across species, whereas the bottom surface (lower) is poorly conserved. B, top view of the gliomedin OLF domain. Residues that are fully conserved in OLF domains are shown as spheres. C, sequence alignment of rat gliomedin (GLDN), sea urchin amassin (AMAS), and various human OLF domains (OLF4, olfactomedin; MYOC, myocilin; LPHN, latrophilin). The fully conserved residues are highlighted in B. The triangles below the alignment highlight some of the most severe mutations in human myocilin (green) and the location of the calcium-coordinating mutation site D380A (blue).
The OLF Domain Is a Monomer Folded into β-Structure in Solution—Gliomedin contains a collagen-like domain N-terminal to its OLF domain. In line with this, EM imaging has shown that gliomedin forms trimers through its collagen-like domain, such that the three OLF domains lie rather close to each other in space (8). To validate the crystal structure, which gives no indications of specific oligomerization, we characterized the OLF domain in solution.

Correct folding of the gliomedin OLF domain in solution was proven by CD spectroscopy, which shows clear hallmarks of β-sheet (Fig. 5A). We further characterized the domain in solution using SAXS, and it behaves like a monomer (Fig. 5B). The SAXS data indicate a radius of gyration ($R_g$) of 2.4 nm and a maximum dimension ($D_{max}$) of 9 nm. These values are larger than those calculated (53) from the crystal structure ($R_g = 1.7$ nm, $D_{max} = 5.3$ nm), which is most likely caused by the N terminus, which is not observed in the crystal structure. The molecular mass, based on the excluded volume of the ab initio model, is 36 kDa, which is very close to that expected for a monomer (32.4 kDa). In three-dimensional modeling, good fits were obtained when the N terminus was implicitly modeled (Fig. 5, B–D). Also the OLF domain of myocilin has been characterized as a monomer in vitro (54, 55).

Structural modeling based on solution scattering data indicates that the construct used in our studies includes an extension to the OLF domain; this most likely represents the disordered N terminus of the protein (Fig. 5, C and D). Although the OLF domain is sufficient to induce Na+ channel clustering, this process requires a pre-clustering of OLF domains (7, 8). Considering the fact that the disordered N terminus of our construct essentially lies at the end of the collagen-like trimerization domain, based on the SAXS model, we calculated the expected dimensions of the C-terminal OLF domain assembly in full-length gliomedin. Assuming the collagen-like domain indeed forms a triple helix, the interaction (top) surfaces of the OLF domains cannot be farther apart from each other than 14 nm (Fig. 5E). Of course, it is possible that they actually are closer to one another in the trimeric molecule.

Comparison with Other OLF Family Members—Within the OLF family, by far the largest body of research has been carried out on myocilin, which is understandable based on its direct involvement in human disease. Dozens of myocilin mutations in glaucoma patients have been described, and the vast majority of these map to the OLF domain (14), highlighting the crucial role of this domain in the physiological function of myocilin.
Crystal Structure of the Olfactomedin Domain

FIGURE 6. Analysis of glaucoma mutations in the myocilin OLF domain. Top (left) and side (right) views are shown to highlight the three-dimensional clustering of known mutations based on a homology model. The Cα atoms of affected residues are shown as spheres. Mutations described as severe (14) are shown in red; all other mutations are shown in green. The side view indicates very few mutations in the secondary structure elements. The orange ellipsoids highlight mutation clustering in the two views.

Using the gliomedin OLF domain crystal structure, a homology model was made for myocilin, which allowed mapping of the OLF domain glaucoma mutations onto the three-dimensional structure (Fig. 6). A number of observations on these mutations can be made. The first striking observation is that despite the β-propeller fold and reports on misfolding and/or abnormal processing of myocilin mutants (56–60), very few of the mutations are actually located in the β-strands; most of them affect loops between the strands. Analysis of mutation location in three-dimensional space further indicates hot-spot regions with highly concentrated mutations; this concerns especially the top face of blades 2–4 and, to some extent, the bottom side of blades 1 and 5 (Fig. 6). The locations of some of the most severe mutations (14) are discussed below.

DISCUSSION

OLF domain proteins are known to play essential roles in development and cell differentiation in both vertebrates and invertebrates; however, their functions are only partially known at the moment, and their high-resolution structures have remained unknown. We have presented the first crystal structure of an OLF domain, which in gliomedin is crucial for correct formation of the nodes of Ranvier in myelinated peripheral nerves. At a more general level, the structure can be used both to understand OLF domain function in different OLF family proteins and to predict and analyze the molecular effects of human disease-causing mutations affecting OLF domains.

In vertebrates, gliomedin is important for the clustering of Na⁺ channels at the nodes of Ranvier (61), a process that is a prerequisite for saltatory propagation of the action potential along axons. On the other hand, the gliomedin homolog anc-122 in Caenorhabditis elegans is involved in neuromuscular signaling (11). Interestingly, the Na⁺ channel-clustering function of gliomedin can also be carried out by the isolated OLF domain (7, 8) as long as the OLF domain has been pre-clustered. Hence, the OLF domain crystal structure presented here directly represents the functional unit of gliomedin in the development of the nodes of Ranvier in the nervous system. Whether the clustering of three OLF domains together by the collagen triple helix is enough for this function or if larger scale aggregates are required in vivo is currently unclear.

A high-resolution structural study including the collagen-like domain N-terminal to the OLF domain would clarify how the three OLF domains are arranged in a gliomedin trimer. EM imaging provided a low-resolution view of this arrangement (8), in which the three OLF domains are close to each other in space, but not bound to each other. This fits our SAXS data and the model of a trimERIC arrangement of monomERIC OLF domains (Fig. 5E). In general, oligomerization via neighboring domains or disulfide bridges is a general feature of OLF domains; for example, myocilin has a coiled-coil dimerization domain instead of the collagen-like domain of gliomedin. Oligomerization of OLF domains through these adjacent regions thus appears to be crucial for their biological activity.

Gliomedin is known to bind to fibronectin type III-like domains of neurofascin and NrCAM (9). These domains contain locally concentrated, negatively charged residues, which could provide a binding site for the positively charged top face of the gliomedin OLF domain. However, the precise binding mode will remain unclear until structural information becomes available for such protein–protein complexes.

Five-bladed β-propeller structures are most often carbohydrate-binding proteins and/or enzymes (49). For gliomedin, apart from its interactions with neurofascin and NrCAM (5), few functional details are known at the molecular level. No enzymatic activity has been reported for OLF domains, and gliomedin is unlikely to harbor one.

Like many other myelin proteins, gliomedin is a target for autoimmune antibodies in peripheral neuropathies (62–64), with implications for multifocal motor neuropathy, chronic inflammatory demyelinating polyneuropathy, and Guillain–Barré syndrome. The exact autoimmune epitopes on gliomedin are not yet known. The crystal structure of the OLF domain will now enable the identification of such epitopes to understand the effects of autoimmune attack on the nodes of Ranvier and the myelin sheath.

Myocilin is by far the best characterized OLF domain-containing protein; however, no published data exist on its three-dimensional structure. The gliomedin OLF domain structure enables us to understand the locations and effects of the plethora of human myocilin mutations causing glaucoma. In three-dimensional space, the human myocilin mutations clearly cluster in defined regions and are largely absent from the secondary structure elements (Fig. 6). This is surprising, as glaucoma mutations in general have been assumed to cause misfolding or abnormal processing of myocilin (56–60). The loops are apparently important for correct OLF domain folding.

A calcium-binding site was reported in myocilin, and it was suggested to involve Asp-380 (52). The glaucoma-causing mutation D380A has been intensively studied with regard to the loss of Ca²⁺-binding affinity upon mutation (52). In the gliomedin structure, we detected a bound cation (most likely sodium) that is coordinated by Asn-423, corresponding in sequence to Asp-380 in myocilin (Figs. 3E and 4C). Thus, Asp-380 in myocilin is most likely located in a similar cavity and possibly chelates a calcium ion that stabilizes the structure. In all other OLF domains apart from gliomedin, this Asp is conserved (Fig. 4C), and calcium binding may be a general feature of the OLF domain, with the exception of gliomedin. Interest-
ingly, calcium is also required for the massive intercellular clotting involving the sea urchin OLF protein amassin (65, 66); it is not known, however, if this reflects a requirement of calcium binding for a functional amassin OLF domain. The second metal-coordinating side chain of gliomedin (Asn-471) is conserved in myocilin as Asn-428. We believe that the sodium-binding site observed in the gliomedin crystal structure spatially corresponds to the calcium-binding site of myocilin. Taking into account that the extracellular Na+ concentration is expected to be higher than that of our buffers, it is likely that the site in gliomedin is also occupied by Na+ in vivo. Furthermore, it is interesting to note that gliomedin is localized close to the nodes of Ranvier, at which Na+ will rush into the axon when a nerve impulse passes, and that it is involved in the clustering of Na+ channels. Previously, Ca2+ binding by myocilin was reported to increase the stability of the OLF domain (52), and the binding of metals could clearly be a common mechanism to increase OLF domain stability. The functional relevance of metal binding by OLF domains clearly requires further study.

The residues corresponding to the myocilin mutations P370L and C433R (responsible for severe glaucoma phenotypes) are located on the loops between β-strands 2d and 3a and between β-strands 4a and 4b, respectively. These residues and the conformation of the loops are probably critical to forming and maintaining a stable β-propeller structure. Furthermore, Cys-433 has been suggested to form a disulfide bridge with Cys-245 in myocilin (67); in gliomedin, such a disulfide is missing. Cys-433 is conserved in all other OLF domains apart from gliomedin. Considering the myocilin homology model, Cys-433 is in close contact with Gly-246, which is the N-terminal residue of the homology model. Thus, Cys-245 would be at a perfect position to form a disulfide bond with Cys-433, and this linkage can be expected to stabilize the assembly of the myocilin OLF domain.

Another human glaucoma mutation is Y437H, which also can induce disease when injected into mice (68). The corresponding residue in gliomedin (Tyr-480) lies in the middle of β-strand 4b, surrounded by many hydrophobic residues. This indicates that Tyr-480 in gliomedin, which indeed is conserved in all OLF domains (Fig. 4C), may be a key residue to stabilize the β-propeller structure.

Of the 20 glycine residues in the myocilin OLF domain, a total of eight have been listed as glaucoma mutation sites (14). Essentially all these residues are located in loops between β-strands in the propeller blades. It is likely that their flexibility and their ability to support tight turns are of overall importance during the correct folding of the myocilin OLF domain. It is also logical to assume that glycine residues in the loops are involved in the folding of other OLF family members.

To conclude, we have presented the crystal structure of the OLF domain from gliomedin, which forms a twisted five-bladed β-propeller. The high-resolution OLF domain structure is an important step toward understanding the functions and mechanisms of OLF domains. It is likely that the tertiary structures of other OLF domains are highly similar to that of gliomedin. Hence, the OLF domain is a unique β-propeller, and OLF-containing proteins can be added to the list of β-propeller proteins involved in human development and disease.

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