Research Article

Mouse Ficolin B Has an Ability to Form Complexes with Mannose-Binding Lectin-Associated Serine Proteases and Activate Complement through the Lectin Pathway

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Ficolins (FCN/Fcn) are a family of proteins comprising a collagen-like and a fibrinogen-like domain [1] the latter binding specifically to N-acetyl compounds such as N-acetylglucosamine (GlcNAc) [2–4]. Three types of ficolin have been identified in humans: L-FCN [1, 5], M-FCN [6, 7], and H-FCN [8]. Transcripts for L-FCN and H-FCN are mainly produced in the liver, and the proteins circulate as serum ficolins, whereas the mRNA for M-FCN is expressed mainly in peripheral monocytes and the protein is present in the serum at low concentrations [9]. Two ficolins have been identified in mice: ficolin A (FcnA) and ficolin B (FcNB) [10, 11]. FcnA mRNA is mainly expressed in Kupffer cells in the liver [12], and the protein is present in serum. FcnB mRNA is mainly expressed in cells of myeloid cell lineage within the bone marrow [12]. The location of FcnB protein is still unclear, although it is reported to localize within the lysosomes of activated macrophages [13].

Our phylogenetic analyses show that FcnB is the murine orthologue of human M-FCN, that FcnA and L-FCN were independently diverged in the murine and primate lineages, respectively, from the ancestral FcnB/M-FCN, and that the H-FCN gene is a pseudogene in the murine lineage [14]. In addition, our previous ontogenetic study showed that the spatial-temporal expression pattern was different for FcnB and FcnA, suggesting that each of the ficolins might have a specific role in the prenatal and postnatal stages [12].

Thus, ficolins are roughly classified into two groups: a serum type (plasma type), which includes L-FCN, H-FCN and FcnA produced mainly in the liver and present in the serum, and present in the circulation as serum lectins [15] and a nonserum type (nonplasma type), which includes M-FCN and FcnB hardly detectable in the serum. The latter group, particularly murine FcnB, has not been studied in detail at the protein level, because of difficulties in identifying and isolating a sufficient amount of the protein. There are no reports regarding the biochemical features of native FcnB.
To date, we have shown that mammalian ficolins, including three human ficolins and mouse FcnA, associate with mannose-binding lectin (MBL)-associated serine proteases (MASPs) and activate the lectin pathway [5, 16–18]. We also reported that recombinant mouse FcnB produced in *Drosophila* S2 cells does not associate with MASP-2 and small MBL-associated protein (sMAP) [18]. Recently, however, it was reported that rat recombinant FcnB associates with MASPs and activates the lectin pathway by binding to PAMPS [19]. In the present study, we carefully examined the biochemical properties of FcnB using both native FcnB isolated from mouse bone marrow fluid, and recombinant mouse FcnB produced in CHO cells. The results show that like rat FcnB, mouse FcnB has the ability to form complexes with MASPs and sMAP.

2. Materials and Methods

2.1. Preparation of FcnB from Mouse Bone Marrow. To avoid complications resulting from the co-presence of FcnA and FcnB, the bone marrow tissue used as the source of FcnB was collected from FcnA-deficient mice generated on a C57BL6 background by gene targeting (manuscript in preparation). The bone marrow fluid and cells were collected as supernatant and precipitate, respectively, by centrifugation (preparation). The bone marrow fluid and cells were subjected to affinity chromatography on a GlcNAc-agarose column. The bound fraction was eluted sequentially with 0.3 M mannose and then with 0.3 M GlcNAc. The recovered eluate was dialyzed against 50 mM Tris-HCl, pH 7.5, containing 0.15 M NaCl and 2.5 mM CaCl2 (TBS-Ca), concentrated, and stored at −80 °C until use. The GlcNAc-eluate was used as a source of native FcnB in further study.

2.2. Preparation of Recombinant FcnB (rFcnB). Full-length mouse FcnB cDNA was constructed in a pIRCMV vector and cotransfected with a pFerH vector encoding a transposase (manuscript in preparation). The bone marrow fluid and cells were collected as supernatant and precipitate, respectively, by centrifugation of the pooled tissue at 10,000 ×g for 10 min. The supernatant was then subjected to affinity chromatography on a GlcNAc-agarose column. The bound fraction was eluted sequentially with 0.3 M mannose and then with 0.3 M GlcNAc. The recovered eluate was dialyzed against 50 mM Tris-HCl, pH 7.5, containing 0.15 M NaCl and 2.5 mM CaCl2 (TBS-Ca), concentrated, and stored at −80 °C until use. The GlcNAc-eluate was used as a source of native FcnB in further study.

2.3. Preparation of Recombinant Mouse MASP-1, MASP-2, MASP-3, and sMAP. Two recombinant forms of mature mouse MASP-2 were produced in *Drosophila* S2 cells with a histidine (His)-tag as previously described [21]: one comprised the normal sequence with protease activity (rMASP-2a) and the other a mutated sequence (Ser632Ala) with no activity (rMASP-2i). Recombinant mouse MASP-1 (rMASP-1i) was prepared in a His-tagged form using a Baculovirus expression system (Invitrogen, Carlsbad, CA, USA), and is an inactive form harboring a mutated catalytic site (Ser646Ala) [22]. Recombinant mouse MASP-3 (rMASP-3) [23] and sMAP (rsMAP) [21] were prepared as His-tagged forms in *Drosophila* S2 cells. All recombinants (rMASP-1i, -2i, -2a and -3, and rsMAP) were purified by affinity chromatography on Ni-NTA agarose columns (Qiagen Inc., Valencia, CA, USA) followed by elution with imidazole. The recombinant proteins were dialyzed against TBS, concentrated, and stored at −80 °C until use. The protein concentration was determined as described above.

2.4. Western Blotting for FcnB, MBLs, MASPs, and sMAP. SDS-PAGE was performed on 10% polyacrylamide gels under reducing conditions according to the method of Laemmli. After electrophoresis, the proteins were transferred to a polyvinylidene difluoride membrane filter (Millipore, Billerica, MA, USA). The membrane filter was blocked with Blocking One reagent (Nacalai Tesque Inc., Kyoto, Japan) and probed with 500-fold-diluted polyclonal antibodies (Ab) against mouse FcnB and MASP-2/sMAP [18, 21] and 500-fold-diluted monoclonal Abs against MBL-A and MBL-C (clones 8G6 and 16A8, resp., Hycult Biotechnology, Uden, The Netherlands) in 10 mM phosphate buffer, at pH 7.4, containing 137 mM NaCl and 2.7 mM KCl (PBS) containing 0.1% Tween-20 (PBS-T). For the detection of MASP-1 and -3, a monoclonal anti-penta-His-tag Ab (Qiagen) was used as the primary Ab. After washing, the filters were further incubated with either HRP-conjugated secondary Abs or biotinylated secondary Abs (Dako Cytomation, Glostrup, Denmark) followed by an avidin-biotinylated HRP complex (Vector Lab., Burlingame, CA, USA). Finally, the membranes were developed using a chemiluminescent substrate (ECL, Amersham Biosciences, Buckinghamshire, UK). The chemiluminescent image was analyzed using an LAS-3000 (Fuji film, Tokyo, Japan).

2.5. Treatment of FcnB with Endoglycosidases. The N-linked carbohydrates expressed on FcnB were removed by treatment with endoglycosidase F (EMD Biosciences Inc., La Jolla, CA, USA) as previously described [24]. Selective removal of O-linked glycans was achieved by treatment with 0.1 U neuraminidase (Wako Pure Chemicals, Osaka, Japan) and 20 mU endo-α-N-acetylgalactosaminidase (Seikagaku Co., Tokyo, Japan) at 37 °C for 16 h.
2.6. Gel Filtration Chromatography of FcnB. To estimate the size distribution of oligomeric FcnB, the FcnB preparations were subjected to gel filtration chromatography using a Superose 6 10/300GL column equilibrated with PBS and connected to an AKTA purifier system (Amersham Biosciences, Uppsala, Sweden). An aliquot of each recovered fraction (0.5 mL/fraction) was assessed for FcnB by western blotting under reducing conditions.

2.7. Complex Formation of rFcnB with rMASPs and rsMAP. rFcnB was incubated with rMASPs and rsMAP at a molar ratio of 3:1:8 (rFcnB: rMASPs: rsMAP) overnight at 4°C in TBS containing 2.5 mM CaCl₂, 3% BSA, and 0.05% Tween-20 as previously described [18]. The above molar ratio was chosen by reference to the concentrations of FcnA, MASP-2, and sMAP in the mouse serum [18]. The mixture was further incubated with a GlcNAc-agarose slurry (50%, 40 μL) at 4°C for 3 hr to pull down rFcnB, and the bound fraction was eluted with 0.3 M GlcNAc. The eluate was dialyzed against TBS-Ca and the final sample subjected to western blotting and a C4-deposition assay. For western blotting, rMASP-2i was used as a source of MASP-2 to ensure clear results, since it is known that rMASP-2a is converted, in part, into its active form, comprising the heavy and light chains connected via a disulfide bond, during purification. For the C4-deposition assay, MASP-2a was used as the source of MASP-2 instead of rMASP-2i. Similar autoactivation is also seen with rMASP-1; therefore, rMASP-1i was used as the source of MASP-1 for western blotting to detect complex formation with rFcnB.

2.8. C4-Deposition Assay. C4-deposition activity was determined by ELISA as previously described [18]. Briefly, the GlcNAc eluates prepared from bone marrow fluid or the rFcnB/rMASP-2a/rsMAP complexes, were incubated in 100 μL of TBS-Ca at 37°C for 10 min in a GlcNAc-BSA-coated microtiter plate. The plate was then incubated with human C4 on ice for 30 min, followed by washing with PBS-T. The C4b generated on the plate was detected with an HRP-conjugated sheep anti-human C4b Ab (Biogenesis, Poole, UK) and color developed using TMB (KPL Co., Gaithersburg, MD, USA) and H₂O₂ as substrates. After termination of the reaction with 0.5 M H₃PO₄, the plates were read at 450 nm in a Multimode detector DTX880 (Beckman Coulter Inc., Brea, CA, USA).

3. Results

To detect the FcnB protein in the bone marrow, the tissue supernatants and precipitates were subjected to western blotting. As shown in Figure 1(a), a 38 kDa band was observed in the supernatant under reducing conditions, suggesting that FcnB is secreted into the mouse bone marrow fluid as a soluble protein. FcnB was also detected as a 37 kDa band at high levels in the precipitate. This suggests that FcnB in bone marrow cells is slightly small due to incomplete processing prior to secretion. When FcnB in the supernatant was treated with endoglycosidase F, its molecular weight reduced from 38 to 34 kDa, whereas treatment with endo-α-N-acetylgalactosaminidase resulted in either no or a smaller reduction in molecular weight (Figure 1(a)). To determine the size distribution of oligomeric FcnB, the supernatant was subjected to gel chromatography. FcnB was recovered from fractions corresponding to the elution positions of marker proteins ranging from 100 to >1000 kDa with a peak around 600 kDa (Figure 1(b)), indicating a heterogeneous structure composed mainly of 12–18-mers.

Next, the bone marrow supernatant was subjected to GlcNAc-agarose affinity chromatography to purify FcnB. As shown in Figure 2(a), FcnB was recovered in the GlcNAc eluate, whereas MBL-A and MBL-C (MBLs) were recovered in the mannose eluate. The mannose eluate included significant amounts of the MASP-2 pro-enzyme, MASP-2 heavy chain, and sMAP. This suggests that MBLs are present in the bone marrow fluid as complexes with MASP-2 and sMAP. Interestingly, trace amounts of MASP-2 and sMAP were also detected in the GlcNAc eluate, suggesting that at least a part of FcnB also exists in complex with MASP-2 and sMAP. This GlcNAc eluate showed C4-deposition on GlcNAc-coated microplates (Figure 2(b)), although the level was very low compared with that of the mannose eluate. This activity was significantly decreased by passage of the eluate through anti-FcnB Ab-coupled Sepharose 4B. These results suggest that the FcnB/MASPs complexes can activate complement component C4 through the lectin pathway.

To confirm the above results, rFcnB was produced in CHO cells and purified by GlcNAc-agarose chromatography. Western blotting showed that rFcnB consisted of a monomer with a molecular weight of 37 kDa; slightly smaller
Figure 2: GlcNAc-agarose chromatography of bone marrow fluids from FcnA−/− mice. (a) Western blot of FcnB, MBL-A, MBL-C, and MASP-2/sMAP in the mannose (M) and GlcNAc (G) eluates from GlcNAc-agarose chromatography under reducing conditions. Right panel: 90 kDa, 58 kDa, and 23 kDa bands represent the proenzyme form of MASP-2, the heavy chain of MASP-2, and sMAP, respectively. For each sample, 60 μL equivalent to the original volume of bone marrow tissue was loaded per lane. (b) C4-deposition activity of the GlcNAc eluate on GlcNAc-BSA-coated microplates. Before assessment, the GlcNAc eluate was passed through anti-FcnB Ab-coupled Sepharose 4B (Ab+) or not (Ab−). The activity of a 30 μL sample equivalent to the original bone marrow tissue was determined in quadruplicate (mean ± SD). Inset: western blot of FcnB in the eluates used for C4 deposition.

Figure 3: Structural characterization of rFcNb. (a) Left panel: Western blot of rFcNb and rFcNbS2-1 treated with endoglycosidase F (endoF). Right panel: western blot of rFcNb and rFcNbS2-1 treated with neuraminidase (neu) or neuraminidase plus endo-α-N-acetylgalactosaminidase (o-gly). Western blotting was performed under reducing conditions. —, not treated. (b) Gel chromatography of rFcNb (upper panel) and rFcNbS2-1 (lower panel). rFcNb or rFcNbS2-1 (400 μL; 5–10 μg) was applied to a Superose 6 column (1 cm φ × 30 cm) and fractionated into 0.5 mL/fractions. An aliquot of each fraction was subjected to western blotting for FcnB under reducing conditions.

than native FcnB, but larger than rFcNbS2-1 (33–35 kDa) (Figure 3(a)). Upon treatment with endoglycosidase F, the molecular weights of rFcNb and rFcNbS2-1 were reduced to 33 kDa and 31–33 kDa, respectively. Treatment of rFcNb with endo-α-N-acetylgalactosaminidase resulted in a slight reduction in the molecular weight to 36 kDa, while treatment of rFcNbS2-1 had no effect. The N-terminal amino acid sequence of rFcNb was T20CPELKV, indicating that the preceding 19 amino acids were removed as a signal peptide by the host CHO cells. The N-terminal sequences of the 35 kDa and 33 kDa bands of rFcNbS2-1 were RSPWPGVFV15HAAG and A18GTCPEL, respectively, indicating that the 35 kDa band corresponded to our designed rFcNbS2-1 product containing the eight plasmid-derived amino acids (underlined) at the N-terminal [18], and the 33 kDa band was another rFcNbS2-1 product with a different N-terminal, which was processed by Drosophila S2 cells.

Oligomeric rFcNb was subjected to gel chromatography to determine its size distribution. It was found that, like native FcnB, the main rFcNb species was eluted in the
FIGURE 4: Complement activation by rFcnB/rMASP-2/rsMAP complexes. Upper panel: western blot of FcnB and MASP-2/sMAP. After incubation of the recombinant proteins as shown in the table, the generated complexes were pulled down by GlcNAc-agarose and subjected to western blotting. Lower panel: C4-deposition activity of the similar pull-down samples on GlcNAc-BSA-coated microplates. The samples used for C4 deposition were prepared the same as those for western blotting except for employment of rMASP-2a instead of rMASP-2i. The activity was determined in quadruplicate (mean ± SD).

range corresponding to 100 to >1000 kDa with a peak around 600 kDa (Figure 3(b)). A minor band was observed at 33 kDa in the rFcnB preparation eluted between 100 and 200 kDa. These results indicate that the rFcnB preparation contains a major 12–18-mer made up of 37 kDa monomers, and a minor 3–6-mer made up of 33 kDa monomers. Gel chromatography of rFcnBs2-1 showed that this protein ranged from 100 to 300 kDa, suggesting that it is a 3–9-mer composed of 33–35 kDa monomers.

To confirm the interaction between rFcnB and MASP and sMAP, rFcnB was incubated with rMASP-2i and rsMAP and then subjected to FcnB pull down using GlcNAc-agarose. As shown in Figure 4, rMASP-2i and rsMAP were coprecipitated only in the presence of rFcnB. Coincubation of rMASP-2i and rsMAP resulted in reduced binding to rFcnB compared with incubation with each alone, suggesting their competitive bindings to rFcnB. The rFcnB/rMASP-2a complex activated C4 on GlcNAc-coated microplates. This C4-deposition activity was inhibited in part by rsMAP. In addition, rFcnB bound to rMASP-1i and rMASP-3, and this binding was partially inhibited by coincubation with rsMAP (Figures 5(a) and 5(b)), suggesting that FcnB associates with all types of MASP and sMAP in a similar manner. No activation of rMASP-3 was observed under these experimental conditions, even when it was complexed with rFcnB on GlcNAc.

4. Discussion

The present study clearly indicates that, like rat recombinant FcnB, both native and recombinant forms of mouse FcnB associate with MASP and sMAP. It also demonstrates that FcnB/MASP/sMAP complexes activate C4 on immobilized GlcNAc. Taken together with the results of Girija et al. [19], these results suggest that at least a part of murine FcnB essentially executes its function through the lectin pathway.

In the present study, it was observed that the monomer size of FcnBs2-1 was smaller than that of the native FcnB and rFcnB proteins, largely due to the N-linked carbohydrate content. It was also found that FcnBs2-1 formed smaller oligomers (3–9-mers), in contrast to the highly oligomeric forms of the native FcnB and rFcnB (12–18-mers). These results simply suggest that the processing of proteins in insect cells is different from that in mammalian cells. To confirm this in the present study, we prepared a third form of recombinant mouse FcnB, termed rFcnBs2-2, in Drosophila S2 cells. The N-terminal residue of rFcnBs2-2 was adjusted to Thr20 as same as that in rFcnB, which was performed by ligation of a FcnB cDNA containing an extra six bases into a pMT/Bip/V5-His A vector. The generated FcnBs2-2 showed a molecular weight of 31 kDa under reducing conditions, and treatment with endoglycosidase F reduced this to 30 kDa (data not shown). It was also found that rFcnBs2-2 was less oligomeric, existing mainly as 3–6-mers, and that it failed to associate with rMASP-2i or rsMAP (data not shown). These results clearly indicate that the processing of recombinant mouse FcnB in Drosophila S2 cells is different from that in mammalian cells.

An interesting result was observed in gel chromatography of the rFcnB preparation, which contained a major and highly oligomeric species comprising 37 kDa monomers and a minor and poorly oligomeric species comprising 33 kDa monomers (Figure 3(b)). In a preliminary study, we observed that culture of CHO cells in the presence of tunicamycin, an inhibitor of N-linked glycosylation, resulted in the preferential production of rFcnB comprising 33 kDa monomers, which was also less oligomeric (3–6-mers) (data not shown). These results suggest that full N-linked glycosylation of FcnB is essential for high level of oligomerization.

We recently observed that Drosophila S2 cells produced trimers of human H-FCN, while CHO cells produced highly oligomeric H-FCN (~18-mers), which were structurally similar to native H-FCN in human serum (data not shown). However, Drosophila S2 cells do not always produce less oligomeric ficolin. For example, this cell line successfully produced highly oligomeric forms of mouse FcnA and human M-FCN (~600 kDa) [17, 18]. At present, the reason why ficolin molecules are processed differently in Drosophila S2 cells it is not known. One possibility might be a small difference among these ficolins in the amino acid sequence that forms the N-linked glycosylation site, for example, the surrounding sequence around Asn-X-Ser/Thr.
rsMAP was performed using an anti-penta-His tag Ab.

In the table, samples pulled down with GlcNAc agarose were subjected to western blotting. Western blotting for rMASP-1i, rMASP-3, and technical assistance. This work was supported in part by grants from the Ministry of Education, Culture, Sports, Science and Technology of Japan.

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