Effects of MASP-1 of the Complement System on Activation of Coagulation Factors and Plasma Clot Formation

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Abstract

Background: Numerous interactions between the coagulation and complement systems have been shown. Recently, links between coagulation and mannose-binding lectin-associated serine protease-1 (MASP-1) of the complement lectin pathway have been proposed. Our aim was to investigate MASP-1 activation of factor XIII (FXIII), fibrinogen, prothrombin, and thrombin-activatable fibrinolysis inhibitor (TAFI) in plasma-based systems, and to analyse effects of MASP-1 on plasma clot formation, structure and lysis.

Methodology/Principal Findings: We used a FXIII incorporation assay and specific assays to measure the activation products prothrombin fragment F1+2, fibrinopeptide A (FPA), and activated TAFI (TAFIa). Clot formation and lysis were assessed by turbidimetric assay. Crot structure was studied by scanning electron microscopy. MASP-1 activated FXIII and, contrary to thrombin, induced FXIII activity faster in the Val34 than the Leu34 variant. MASP-1-dependent generation of F1+2, FPA and TAFIa showed a dose-dependent response in normal citrated plasma (NCP), albeit MASP-1 was much less efficient than FXa or thrombin. MASP-1 activation of prothrombin and TAFI cleavage were confirmed in purified systems. No FPA generation was observed in prothrombin-depleted plasma. MASP-1 induced clot formation in NCP, affected clot structure, and prolonged clot lysis.

Conclusions/Significance: We show that MASP-1 interacts with plasma clot formation on different levels and influences fibrin structure. Although MASP-1-induced fibrin formation is thrombin-dependent, MASP-1 directly activates prothrombin, FXIII and TAFI. We suggest that MASP-1, in concerted action with other complement and coagulation proteins, may play a role in fibrin clot formation.

Introduction

The coagulation and complement systems are activated following external injury to protect the host from blood loss and infections. The simultaneous activation of coagulation and inflammatory processes after injury is a phylogenetically ancient adaptive response that can be traced back to early eukaryotic evolution [1]. A number of recent studies show direct interactions between the two systems [2,3], among them are links between coagulation factors and mannan (or mannose) -binding lectin (MBL) associated serine proteases (MASPs) of the complement lectin pathway.

The lectin pathway of the complement system is activated by binding of the target recognition molecules MBL or ficolins to carbohydrates or N-acetylated groups, respectively, on the surface of cells or microorganisms. MBL and ficolins circulate in complexes with MASPs which autoactivate upon binding of MBL/ficolins to their target structures. Three MASPs and two related proteins are present in human plasma, arising from two genes by alternative splicing: Mannose-binding lectin-associated serine protease-1 (MASP-1) and its alternatively-spliced variants MASP-3 and MAp44, and MASP-2 and its alternatively-spliced variant MAp19. MAp44 and MAp19 contain no serine protease domain and hence lack enzymatic activity [4,5]. Average plasma concentrations of MASP-1 have been estimated at 6 mg/ml [6] and 11 mg/ml [7].

Upon activation, MASP-2 cleaves both C4 and C2 and thus induces further complement activation by generating the lectin/classical pathway C3 convertase C4b2b. Furthermore, MASP-2 cleaves prothrombin to generate active thrombin [8]. In contrast,
the physiological roles of MASP-1 and MASP-3 remain subject of research although potential substrates have been identified. MASP-3 does not activate either C4 or C2. MASP-1 cleaves C2 but not C4 and therefore is not capable of generating C5 convertase alone [9], as confirmed by a study in MASP-2 knockout mice [10]. Direct activation of C3 by MASP-1 occurs with very low catalytic efficiency and is not of physiological relevance [4,5]. However, MASP-1 has been suggested to act synergistically with MASP-2 to produce C3 convertase via C2 cleavage [11], and MASP-1 may even activate MASP-2 [12,13]. A new role for MASP-1 in the alternative complement pathway has been proposed, by directly activating complement factor D [14,15]. MASP-1 may also exert proinflammatory effects through releasing bradykinin from high-molecular weight kininogen [16].

Experiments with synthetic peptides and structure-function analyses based on its crystal structure have revealed that MASP-1 has a much broader substrate specificity than MASP-2 and a thrombin-like activity with similar substrate specificity [9,17,18]. Indeed, MASP-1 is able to cleave the main substrates for thrombin, fibrinogen and factor XIII (FXIII), and activate endogenous protease-activated receptor 4, albeit with a lower catalytic efficiency compared with thrombin [4,19,20]. Cleavage of the FXIII A-subunit and the fibrinogen β-chain by MASP-1 occur at the same site as proteolysis by thrombin, whereas the cleavage site in the fibrinogen α-chain is different [19], indicating potential differences in the mechanisms of fibrin formation by thrombin and MASP-1. Activation of MASP-1 in complex with L-ficolin or MBL was also associated with generation of a fibrin clot [21]. First evidence for an in vivo role of MASP-1 in coagulation has been recently documented in a study demonstrating that mice lacking MBL or MASP-1 have a prolonged bleeding time following tail tip excision [22]. Furthermore, it has been recently demonstrated in a mouse model of FeCl3-induced intra-arterial thrombogenesis, that MASP-1 plays a key role in thrombus formation in vivo [23].

So far, studies on MASP-1 interactions with clotting factors mainly employed purified systems and cleavage/activation products of fibrinogen and FXIII were observed by Western Blotting and HPLC, but the effects of MASP-1 on clot formation and fibrinolysis in plasma or the characteristics of the clot were not directly assessed. Fibrin clot structure can determine predisposition to thrombotic events [24,25], which is related, at least in part, to fibrinolysis efficiency as clots with denser structure and thinner fibers are generally more difficult to lyse [26]. The relationship between clot structure and lysis is in turn related to incorporation of different plasma proteins into the fibrin network [27]. A common genetic variant of FXIII, Val34Leu, influences FXIII activation and cross-linking efficiency leading to altered fibrin structure [28,29]. Because of its high allele frequency and impact on thrombotic risk, the FXIII Val34Leu polymorphism is of major interest when studying fibrin formation and structure in the context of vascular diseases, but it has so far not been taken into account in studies investigating MASP-1-induced FXIII activation. Finally, given the thrombin-like activity of MASP-1, it is possible that MASP-1 also interacts with another thrombin substrate, thrombin-activatable fibrinolysis inhibitor (TAFI), which has not been studied before.

Therefore, the objective of this work was to investigate the effect of MASP-1 on activation of thrombin-dependent coagulation factors and fibrin clot formation in plasma environment. The specific aims were to 1) investigate MASP-1-induced plasma FXIII activation and analyse the effect of FXIII Val34Leu polymorphism, 2) investigate fibrinogen cleavage and activation of prothrombin and TAFI by MASP-1, and 3) analyse MASP-1-induced plasma fibrin clot formation, structure and lysis by turbidimetric assays and scanning electron microscopy.

**Methods**

**Recombinant MASP-1**

A recombinant mutant R304Q MASP-1 catalytic fragment, prepared as described earlier [9,18], was used in this study. It has a molecular weight of 45.5 kDa and consists of the three C-terminal domains complement control protein 1 and 2 followed by the serine protease domain (CCP1-CCP2-SP). The mutant is resistant to autolysis and preserves its enzymatic activity during prolonged incubation (estimated half-life of several hours in plasma at 37°C), otherwise it has the same enzymatic properties as the wild-type enzyme. To simplify we use the term MASP-1 for the R304Q rMASP-1 catalytic fragment.

**FXIII activation**

FXIII activation was assessed using an incorporation assay [28]. Briefly, a microtiter plate was coated with 100 µl/well of fibrinogen (Sigma-Aldrich, Switzerland) at 40 µg/ml (0.1 µM) in tris-buffered saline (TBS; 40 mM Tris, 140 mM NaCl, pH 7.4). After blocking with 200 µl of 1% bovine serum albumin (BSA), plasma diluted 1/10 as FXIII source or purified FXIII was added followed by the activation mix to a final reaction volume of 100 µl. The activation mix contained 5-biotinamidopentylamine (EZ-Link® Pentylamine-Biotin, Thermo Scientific, USA), dithiothreitol (Sigma-Aldrich), and CaCl2, at final concentrations of 0.3 mM, 5 mM, 10 mM, respectively, and human thrombin (Calbiochem, Merck Biosciences, Switzerland) or MASP-1 at various concentrations (as indicated in the results section). The reaction was incubated at 37°C and stopped at different time points by addition of 200 mM EDTA. The biotinylpentylamine incorporated into fibrin by FXIIIa was detected by streptavidine-alkaline phosphatase (final concentration 1 µg/ml; Sigma-Aldrich), developed with p-nitrophenylphosphate as substrate, and read at 405 nm, with 620 nm as reference wavelength, on a microplate reader.

Besides normal citrated plasma (NCP; local plasma pool), we used FXIII-depleted plasma (FXIII-DP; Kordia, The Netherlands), and prothrombin-depleted plasma (PT-DP; American Diagnostica, USA) to show effects independent from thrombin generation. In some experiments, 1 µM C1 esterase inhibitor from human plasma (C1-Inh; Sigma-Aldrich) was used. Recombinant FXIII-A Val34 and FXIII-A Leu34 variants were obtained from Zedira (Germany).

**Fibrin cross-linking by FXIIIa**

Fibrin cross-linking analysis by SDs-PAGE was performed to confirm that fibrin cross-linking occurs in presence of MASP-1. We incubated 100 µl citrated plasma (normal citrated plasma NCP or FXIII-depleted plasma FXIII-DP) with 50 µl CaCl2 (final concentration 5 mM) and 50 µl thrombin (final concentration 0.075 U/ml) or MASP-1 (final concentration 5 µg/ml) at 37°C for 45 min. To two FXIII-DP samples purified FXIII (final concentration 10 µg/ml) was added. After incubation, samples were spun down at 10’000 g for 10 min, the supernatants were removed and the clots washed twice in 1 ml TBS. After the last wash, 800 µl Laemmli sample buffer (Bio-Rad Laboratories, Hercules, USA) with β-mercaptoethanol was added and the samples were boiled at 95°C for 15 min to solubilize the clots. Purified fibrinogen (Calbiochem; final concentration 0.2 µg/ml) was used as control. Twenty microliters were loaded onto a 7.5% Tris-Glycine Mini-Protean® TGX™ gel (Bio-Rad). The gel was stained with Bio-Safe™ Coomassie (Bio-Rad).
Fibrinogen cleavage and activation of prothrombin and TAFI

Cleavage of fibrinogen and activation of prothrombin were studied in NCP and PT-DP. Plasma samples (400 μl) were incubated at 37°C for 30 min with 100 μl of either TBS buffer only (non-activated, negative control), human thrombin (Calbiochem) at a final concentration of 0.1 U/ml (2.8 nM), human FXa (Hyphen BioMed, Neuville-sur-Oise, France) at a final concentration of 0.1 μg/ml (2.2 nM) with 6 mM CaCl2, or MASP-1 at final concentrations of 20 μg/ml (440 nM), 10 μg/ml (220 nM), 5 μg/ml (110 nM), 2.5 μg/ml (55 nM). Samples were then spun down at 10'000 g for 10 min, and the supernatant was snap-frozen in aliquots. For confirmation of prothrombin cleavage in a purified system, 100 μl of human prothrombin (Hyphen BioMed) at 100 μg/ml (1.4 μM) were incubated at 37°C with FXa at a final concentration of 0.1 μg/ml (2.2 nM) and 2.5 mM CaCl2 or MASP-1 at final concentrations of 20 μg/ml (440 nM), 10 μg/ml (220 nM), 5 μg/ml (110 nM), 2.5 μg/ml (55 nM). At 5, 10, 15, and 30 min samples were snap-frozen. F1+2 was measured using the Enzygnost® H F1+2 ELISA kit (Siemens Healthcare, Marburg, Germany). FPA was determined with the Zymutest FPA ELISA kit (Hyphen BioMed).

TAFI activation in plasma was analysed with the Actichrome® H TAFI activity kit (American Diagnostica Inc, Stamford CT, USA). NCP and PT-DP, diluted 1/25, were incubated for 20 min with an activation mix containing either thrombin-thrombomodulin complex (final thrombin concentration 80–100 nM) or MASP-1 at final concentrations of 50 μg/ml (1100 nM), 10 μg/ml (220 nM), or 2 μg/ml (44 nM). Non-activated plasma was assayed in parallel. After stopping the activation reaction, activated TAFIa was detected with a highly specific TAFIa substrate. The concentration of TAFIa was calculated as the difference between activated and non-activated plasma samples.

TAFI cleavage by MASP-1 was verified using SDS-PAGE. To investigate the dose-dependent effect of MASP-1 on TAFI, 200 μg/ml (3.45 μM) of purified TAFI (Enzyme Research Laboratories, South Bend, USA) were incubated for 30 min at 37°C with MASP-1 at final concentrations of 100 μg/ml (2.2 μM), 200 μg/ml (4.4 μM), 400 μg/ml (8.8 μM), or 800 μg/ml (17.6 μM), diluted in TBS (40 mM Tris, 140 mM NaCl, pH 7.4). To investigate the time-dependent effect of MASP-1 on TAFI, 200 μg/ml (3.45 μM) of purified TAFI were incubated for 30 min, 60 min, and 120 min at 37°C without and with 400 μg/ml (8.8 μM) MASP-1. To confirm MASP-1-induced TAFI cleavage at physiological concentrations and compare it to thrombin/thrombomodulin-induced TAFI cleavage, the following experiment was performed: In a reaction volume of 50 μl, we incubated 15 μg/ml TAFI with either 15 μg/ml rMASP-1 or 1.7 U/ml (50 nM) thrombin and 3 μg/ml (58 nM) thrombomodulin (Abcam, Cambridge, UK) at 37°C for 1 h. This was similar to a protocol by Bajzar et al. [30]. After the incubation steps, reactions were stopped by adding Laemmli sample buffer (Bio-Rad Laboratories, Hercules, USA) containing β-mercaptoethanol, followed by boiling at 95°C for 10 min. Samples were run on Any kD™ Mini-Protean® TGX™ gels (Bio-Rad) and gels were stained with Bio-Safe™ Coomassie.

Turbidimetric clot formation and lysis assay

Plasma clot formation and lysis was measured using a method adapted from Carter et al. [31]. Briefly, 25 μl of citrated plasma...
and 75 µl of TBS (50 mM Tris, 100 mM NaCl, pH 7.4) per well were added to a 96-well microtiter plate (Greiner bio-one, Germany). In the lysis assay, the 75 µl of TBS contained tPA (final concentration 83 ng/ml (1.2 nM)) (Technoclone, Austria). Upon addition of 50 µl of activation mix containing MASP-1 at various concentrations (see results section) or 0.03 U/ml (0.8 nM) (final concentration) human thrombin (Calbiochem) and 2.5 mM (final concentration) CaCl₂ in TBS, the plate was placed into a microplate reader and optical density at 340 nm was recorded at 37°C every 30 sec for up to 2.5 hours. We analyzed the parameters lag time (time between addition of activation mix and exponential increase in absorbance), maximum absorbance (corrected for baseline absorbance), and lysis time (time between maximum absorbance and return to baseline absorbance). The turbidimetric assays were performed in NCP and PT-DP (to exclude thrombin action).

**Scanning electron microscopy of plasma clots**

Forty-five microliters of normal citrated plasma diluted 1/2 with TBS were clotted upon the addition of 5 µl activation mix containing CaCl₂ (final concentration 5 mM) and either thrombin (final concentration 0.88 U/ml (24 nM)) or different concentrations of MASP-1 (final concentrations 1 µg/ml (22 nM), 5 µg/ml (110 nM) or 10 µg/ml (220 nM)). As described earlier [27], clots were made in specially devised small perforated plastic vessels, incubated in a moist chamber for 2 hours followed by washing with sodium cacodylate buffer and subsequently fixed for 30 mins in 2% glutaraldehyde. Clots were recovered and further processed.

![Figure 2. Activation of recombinant FXIII Val34 and Leu34 variants by MASP-1.](image)

(A) In a purified system, thrombin activated the Leu34 variant faster than the Val34 variant, whereas MASP-1 activated the Val34 variant faster than the Leu34 variant. (B) When added to FXIII-depleted plasma the same results in regard to Val34Leu variants were observed, but MASP-1-induced FXIII activation was less efficient compared to thrombin.

Data points represent mean values from two experiments with error bars representing standard deviation.

doi:10.1371/journal.pone.0035690.g002
by a stepwise dehydration with acetone gradient and sputter coated with platinum palladium. Samples were viewed and photographed using a field emission gun, environmental scanning electron microscope (FEI, Quanta 200F, Hillsboro, USA). Ten images from different areas at three different magnifications (5000×, 10000×, 30000×) were taken per clot and visually examined by several investigators blinded to the clotting agent. Fiber diameters were measured with image analysis software package ImageJ 1.44p [Wayne Rasband, National Institutes of Health, USA, http://imagej.nih.gov/ij/]. Ten fibers per image, corresponding to 100 fibers per clot, were measured. Fiber diameter data were normally distributed and compared between the four groups of clots (thrombin, MASP-1 10000 g/ml, MASP-1 6000 g/ml, MASP-1 30000 g/ml) by Oneway ANOVA with post-hoc Bonferroni analysis (SPSS Statistics software, version 17.0.0).

Summary of statistical methods
Experiments were performed several times, as described for the individual experiments, and results are shown as mean and standard deviation (SD) unless stated otherwise. In figures, data points represent mean values with error bars representing SD. SPSS Statistics software, version 17.0.0 (SPSS Inc.) was used for statistical analyses. Associations between MASP-1 concentration and parameters of the turbidometric clot formation and lysis assay, i.e. lag time, maximum absorbance, and lysis time, were assessed using bivariate correlation analysis and expressed as Pearson correlation coefficient. Data on fibrin fiber diameters obtained from SEM of thrombin and MASP-1 clots were tested for normal distribution using the Kolmogorov-Smirnov test. Fiber diameters from SEM of thrombin and MASP-1 clots were tested for normal distribution using the Kolmogorov-Smirnov test. Fiber diameters were compared between the four groups (thrombin, MASP-1 10000 g/ml, MASP-1 6000 g/ml, MASP-1 30000 g/ml) by Oneway ANOVA with post-hoc Bonferroni analysis (SPSS Statistics software, version 17.0.0).

Results

FXIII activation
In a purified system with 20 μg/ml (62 nM) of FXIII, MASP-1 induced FXIII-dependent biotin incorporation in a dose-dependent manner, whereas no incorporation was observed without FXIII or in the presence of 100 μg/ml (1 μM) of C1 inhibitor (Fig. 1a). In normal plasma (final dilution 1/25), MASP-1 induced FXIII-dependent biotin incorporation in a time- and dose-dependent manner (Fig. 1b, 1c). In the purified system, MASP-1-induced FXIII activation was evident at 5 min, whereas in plasma environment no effect was observed until 40 min, in contrast to thrombin which showed an early effect. No incorporation was observed in FXIII-depleted plasma (data not shown), whereas MASP-1 also induced FXIII-dependent biotin incorporation in prothrombin-depleted plasma (Fig. 1d), albeit less efficiently compared with normal plasma (Fig. 1b).

When recombinant FXIII A-subunit Val34 and Leu34 variants were tested in TBS containing 1% BSA or in FXIII-depleted plasma, thrombin induced FXIII activity faster in the Leu34 variant compared with the Val34 variant, whereas the results were the opposite with MASP-1 which induced FXIII activity faster in the Val34 variant than the Leu34 variant (Fig. 2a, 2b). In the purified system, activation of FXIII Leu34 by thrombin or MASP-1 was similar, whereas FXIII Val34 activation showed large differences, with far superior activation achieved with MASP-1. In FXIII-DP, MASP-1 was again less efficient at activating FXIII with both Val34 and Leu34 variant demonstrating higher activation by thrombin, in contrast to purified data. However, Val34 variant was still more efficiently activated by MASP-1 compared with Leu, in contrast to thrombin.

Fibrin cross-linking by FXIIIa
Fig. 3 shows fibrin cross-linking by FXIIIa. Upon activation with thrombin and Ca2+ (lane 3), the fibrinogen Aα-chain (labelled α) was cleaved and partly formed high-molecular-weight αn polymers, the fibrinogen ββ-chain (labelled β) was cleaved, and the fibrinogen γγ-chain (labelled γ) completely disappeared to form γ-γ dimers. When NCP was incubated with MASP-1 (lane 4), the same cross-linking pattern appeared. No cross-linking was seen in FXIII-DP (lanes 5 and 6), but could be restored by addition of FXIII (lanes 7 and 8) confirming that this process was depending on FXIII.

MASP-1 and Plasma Clot Formation
When MASP-1 was added to NCP, a dose-dependent generation of the prothrombin activation product, prothrombin fragment F1+2, was observed, although it was far less efficient than FXa (Table 1, Fig. 4a). We confirmed prothrombin cleavage by MASP-1 in a purified system which showed a dose- and time-response (Fig. 4b). In NCP, MASP-1 concentration correlated with generation of the fibrinogen cleavage product FPA, although it was again far less efficient than thrombin (Table 1, Fig. 4c). Using PT-DP, thrombin had similar effects on FPA generation compared with NCP, whereas MASP-1 showed only negligible FPA generation and failed to demonstrate a dose-response effect (Table 1, Fig. 4c). This suggests that in a plasma system, MASP-1-induced FPA cleavage is largely thrombin-dependent and secondary to prothrombin activation by MASP-1.

In addition to prothrombotic factors, MASP-1 is capable of activating the antifibrinolytic protein TAFI. Using NCP, MASP-1 activated TAFI in a dose-dependent manner, although high concentrations were needed to show an effect and this was still
around 10-fold lower than the activation achieved by the physiological TAFI activator thrombin-thrombomodulin complex (Table 1). MASP-1 also activated TAFI in PT-DP indicating that this process is not thrombin-dependent. Cleavage of TAFI by MASP-1 in a dose- (Fig. 5a) and time-dependent (Fig. 5b) manner was confirmed by SDS-PAGE. Again, MASP-1 was less efficient at cleaving TAFI than thrombin/thrombomodulin (Fig. 5c).

Clot formation, structure and lysis using turbidimetric assays

MASP-1 induced clot formation in NCP, but not in PT-DP (not even at MASP-1 concentrations up to 100 µg/ml (2.2 µM)). This is in agreement with our FPA data above and suggests that in plasma environment fibrin formation is not directly induced by MASP-1 in the absence of prothrombin.

MASP-1 had no clear influence on the initiation of clot formation in NCP assessed as lag time. However, addition of MASP-1 resulted in prolongation of lysis time, with maximum prolongation at 1 µg/ml (22 nM) but no additional effect at higher MASP-1 concentrations (Fig. 6a). Furthermore, MASP-1 had a dose-dependent effect on plasma clot maximum absorbance (Fig. 6b). MASP-1 concentrations between 0 and 8 µg/ml (176 nM) showed significant inverse correlation with maximum absorbance (Pearson correlation coefficient $r = 0.964$, $p < 0.001$).

These results suggested that higher MASP-1 concentrations were associated with either thinner fibers or reduction in clot density.

**Fibrin structure by SEM**

Clots prepared with thrombin (Fig. 7d) were characterized by a dense fibrin network with thin fibers with a mean (SD) diameter of 102 (18) nm and small pores. In contrast, clots prepared in presence of MASP-1 (Fig. 7a–c) showed a less dense network with coarser fibers and larger pores. The mean (SD) fiber diameters of clots made with 1 µg/ml (22 nM), 5 µg/ml (110 nM) or 10 µg/ml (220 nM) of MASP-1 were 108 (17) nm, 120 (17) nm, and 134 (22) nm, respectively. Fiber diameters of MASP-1 clots were significantly higher than thrombin clots ($p < 0.001$), except for clots prepared with 1 µg/ml MASP-1 ($p = 0.097$).

**Role for MASP-1 in plasma clot formation**

Taken together, we propose the following role for MASP-1 in plasma clot formation, as summarized in Fig. 8. MASP-1 initiates fibrin clot formation by converting prothrombin to thrombin which then cleaves fibrinogen. MASP-1 activates FXIII independently from thrombin, having a greater effect on Val34 variant compared with Leu34, which is the opposite to thrombin. MASP-1 induces changes in fibrin clot structure that may be related to different plasma protein incorporation into the clot secondary to altered FXIII activation. This, together with direct TAFI

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**Table 1. Activation products of prothrombin, fibrinogen, and TAFI.**

| Activation by            | NCP          | PT-DP         |
|--------------------------|--------------|---------------|
| **Activation by F1+2 (pmol/l)** |              |               |
| FXa 0.1 µg/ml (2.2 nM)   | 136417.0     | Not applicable|
| MASP-1 20 µg/ml (440 nM) | 689.3        | Not applicable|
| MASP-1 10 µg/ml (220 nM) | 461.2        | Not applicable|
| MASP-1 5 µg/ml (110 nM)  | 299.9        | Not applicable|
| MASP-1 2.5 µg/ml (55 nM) | 232.2        | Not applicable|
| No activation            | 136.0        | Not applicable|

| Activation by FPA (nmol/l) |              |               |
|---------------------------|--------------|---------------|
| Thrombin 0.1 U/ml (2.8 nM)| 556.6        | 516.9         |
| MASP-1 20 µg/ml (440 nM)  | 18.7         | 4.2           |
| MASP-1 10 µg/ml (220 nM)  | 12.3         | 4.0           |
| MASP-1 5 µg/ml (110 nM)   | 7.8          | 4.0           |
| MASP-1 2.5 µg/ml (55 nM)  | 4.8          | 4.0           |
| No activation             | <0.4         | 2.4           |

| Activation by TAFIa (nmol/l) |              |               |
|-----------------------------|--------------|---------------|
| Thrombin-thrombomodulin     | 360.3        | 549.1         |
| MASP-1 50 µg/ml (1000 nM)   | 38.9         | 44.0          |
| MASP-1 10 µg/ml (220 nM)    | 7.4          | 10.9          |
| MASP-1 2 µg/ml (44 nM)      | 1.4          | 3.1           |

Activation products were measured in normal citrated plasma (NCP) and prothrombin-depleted plasma (PT-DP). Data are shown as mean ($n = 2$). TAFIa results are corrected for the non-activated samples, therefore no values for the non-activated samples are shown.

doi:10.1371/journal.pone.0035690.t001
Figure 5. Cleavage of TAFI visualized by SDS-PAGE. (A) Dose-dependent effect of MASP-1 on cleavage of TAFI. Lane 1: Molecular weight marker. Lane 2: TAFI 200 µg/ml. Lane 3: TAFI+100 µg/ml MASP-1. Lane 4: TAFI+200 µg/ml MASP-1. Lane 5: TAFI+400 µg/ml MASP-1. Lane 6: TAFI+800 µg/ml MASP-1. (B) Time course of MASP-1 cleavage of TAFI. TAFI (200 µg/ml) was incubated at 37°C without or with MASP-1 (400 µg/ml). When TAFI was incubated without MASP-1, no cleavage occurred. Lane 1: Molecular weight marker. Lane 2: MASP-1 (fresh control). Lane 3: TAFI (fresh control). Lane 4: TAFI incubated for 30 min. Lane 5: TAFI+MASP-1 incubated for 30 min. Lane 6: TAFI incubated for 60 min. Lane 7: TAFI+MASP-1 incubated for 60 min. Lane 8: TAFI incubated for 120 min. Lane 9: TAFI+MASP-1 incubated for 120 min. (C) Physiological concentrations of TAFI (15 µg/ml) were incubated with either physiological concentrations of MASP-1 (15 µg/ml) or 1.7 U/ml (50 nM) thrombin and 3 µg/ml (58 nM) thrombomodulin. Lane 1: Molecular weight marker. Lane 2: TAFI. Lane 3: TAFI+MASP-1. Lane 4: TAFI+thrombin/thrombomodulin. In all three panels, arrows indicate full-length TAFI (58 kDa) and activated TAFIa (35 kDa).

doi:10.1371/journal.pone.0035690.g005

activation by MASP-1, may be responsible for longer lysis time of clots formed following MASP-1 activation.

Discussion

There is increasing interest in and mounting evidence for a role of the complement system in atherosclerosis [32]. In addition, more interactions between complement and coagulation are being discovered which could provide pathophysiological mechanisms linking inflammatory and thrombotic processes. The aim of the present work was to study interactions between MASP-1 and coagulation factors involved in fibrin formation in plasma and investigate whether MASP-1 is able to induce plasma clot formation independently from activation of coagulation pathways. Completely novel aspects of our work compared with earlier studies include the following: 1) experiments were performed in plasma environment, 2) we employed state of the art coagulation tests including sensitive activation tests and a turbidimetric clot formation and lysis test, 3) for the first time we studied the activation of both FXIII Val34 and Leu34 genetic variants by MASP-1, 4) we investigated for the first time MASP-1 effects on TAFI, and 5) we studied for the first time the structure of plasma clots made in presence of MASP-1 using SEM.

We showed that MASP-1 activates FXIII in plasma in a dose-dependent manner and independently from thrombin activity, albeit less efficiently compared to thrombin. MASP-1 specificity was further confirmed by inhibition of FXIII activation in the presence of C1-inhibitor. MASP-1 was slower at activating FXIII in plasma environment, with similar results observed in normal and prothrombin-depleted plasma, than in the purified system. This suggests that plasma contains factors that inhibit MASP-1-induced FXIII activation, but it also shows that MASP-1 activation of FXIII can occur directly and is not only thrombin-dependent. However, in plasma environment most FXIII activation by MASP-1 is probably mediated by thrombin and sufficient FXIIIa is generated to support normal fibrin cross-linking as shown by our cross-linking analysis. On the other hand, it is known that only traces (<1%) of FXIIIa are sufficient for fibrin cross-linking [33], so even in the absence of thrombin the FXIIIa directly generated by MASP-1 would be enough to cross-link any fibrin or other proteins. One striking observation was the difference in MASP-1-induced activation of FXIII genetic variants. The common FXIII Val34Leu polymorphism leads to altered FXIII activation by thrombin [28,29]. The FXIII A-subunit residue at position P(4) from the thrombin cleavage site, corresponding to Val34, is important for recognition and binding of thrombin and hence FXIII activation [34,35]. Due to its higher affinity to thrombin, the Leu34 variant is activated at a faster rate and paradoxically this variant protects from cardiovascular disease in most clinical studies [36]. Studies have struggled to explain this paradox and several hypotheses have been put forward including an interaction with fibrinogen levels and less efficient cross-linking of various proteins into the clot [29,37,38]. In our work, however, we have shown that MASP-1 activated the wild-type Val34 variant faster than the Leu34 variant, both in a purified system and in plasma environment. This suggests that i) MASP-1 activates FXIII in a different manner compared with thrombin possibly due to differences in substrate recognition and ii) interaction between the complement system and FXIII activation in vivo may contribute, at least in part, to the higher prevalence of thrombotic events with FXIII Val34 variant. Further work to elucidate the mechanisms behind this intriguing novel observation remains an area for future studies, including experiments at different fibrinogen concentra-
tions since the protective effect of the FXIII Leu54 variant is strongest at high fibrinogen concentrations.

We also tested MASP-1-induced cleavage/activation of two other major thrombin substrates in plasma, fibrinogen and TAFI. Cleavage of the fibrinogen \( \alpha \)-chain by MASP-1 has been suggested to occur at different sites compared with thrombin [19]. We have therefore tested whether MASP-1 induces the generation of FPA in plasma environment. In NCP, FPA generation correlated with MASP-1 concentration, but the quantities of FPA generated were much lower compared with thrombin. In PT-DP, negligible amounts of FPA were detected which failed to show a MASP-1 dose-response, suggesting that MASP-1-induced release of FPA relies on presence of prothrombin and possibly on thrombin generation. These findings support earlier data that fibrinogen cleavage by MASP-1 does not produce FPA [19] and suggest that direct cleavage of fibrinogen, and FXIII, by MASP-1 is probably of minor importance in plasma. The essential role of prothrombin is supported by our data demonstrating the inability of MASP-1 to induce fibrin clot formation in PT-DP in the turbidimetric assay, and the ability of MASP-1 to cleave prothrombin in a dose-dependent manner. Despite of the apparently better performance of MASP-1 than FXa in cleaving prothrombin in a purified system (Fig. 4b), it must be noted that FXa activity is underestimated in this system due to the lack of its prothrombinase complex cofactor FVa and phospholipids. In plasma (Table 1), prothrombin cleavage by MASP-1, similar to prothrombin cleavage by MASP-2 described by Krarup et al. [8], is much less effective compared to FXa, and would even be less effective when compared with the prothrombinase complex. However, thanks to its efficient amplification system, tiny amounts of generated

Figure 6. Effects of MASP-1 on plasma clot lysis time and maximum absorbance. Clot formation and lysis in normal plasma were investigated using a turbidimetric assay. Increasing MASP-1 concentrations were associated with longer clot lysis time (A) and lower maximum absorbance (B). Data points represent mean values from three experiments with error bars representing standard deviation. doi:10.1371/journal.pone.0035690.g006
thrombin can trigger significant activation of the coagulation cascade. We confirmed MASP-1 cleavage of prothrombin in a purified system, since this reaction had been excluded for MASP-1 earlier [8]. The discrepancy between our results and the study by Krarup et al. [8] may be explained by i) different experimental conditions, ii) difference in concentrations of tested proteins, and iii) different methods to detect prothrombin activation products. Krarup et al. used prothrombin at a final concentration of 110 μg/ml, MASP-1 at 1 μg/ml. We used a similar prothrombin concentration but MASP-1 was used at a wider range of physiological concentrations of 2.5–20 μg/ml. Also, Krarup et al. detected prothrombin cleavage by SDS-PAGE and Coomassie staining. Our method of detecting prothrombin fragment F1+2 by ELISA may be more sensitive.

In contrast to PT-DP, MASP-1 induced clot formation in NCP and an inverse correlation was detected between clot final turbidity and MASP-1 concentrations, indicating this protein modulates the final structure of the clot, which has not been documented before. It is accepted that lower maximum absorbance of clots made using purified systems indicates the formation of thinner fibers, but in plasma clots this is more complex as reduced maximum absorbance may be an indicator of thinner fibrin fibers and/or decreased clot density. MASP-1 has also affected clot structure as assessed by SEM. Clots made in the presence of MASP-1 showed a less dense structure with larger pores and thicker fibers.

Figure 7. Fibrin structure assessed by SEM. Representative images of plasma clots prepared with (A) 1 μg/ml (22 nM) MASP-1, (B) 5 μg/ml (110 nM) MASP-1, (C) 10 μg/ml (220 nM) MASP-1, or (D) 0.88 U/ml (24 nM) of thrombin (magnification 5000×). Clots prepared in presence of MASP-1 showed a less dense structure with larger pores and thicker fibers.

doi:10.1371/journal.pone.0035690.g007

Figure 8. The proposed role of MASP-1 in plasma clot formation. MASP-1 induces fibrin clot formation by converting prothrombin to thrombin which then cleaves fibrinogen. MASP-1 activates FXIII and TAFI independently albeit at a lower efficiency than thrombin. Changes in clot structure and TAFI activation by MASP-1 may be responsible for prolonged lysis time of clots made in the presence of MASP-1. The finer arrows for MASP-1-dependent reactions, as compared with thrombin-dependent reactions, represent the fact that in plasma MASP-1 is less efficient compared with thrombin. The mechanisms underlying the impact of MASP-1 on fibrin structure remain to be elucidated.

doi:10.1371/journal.pone.0035690.g008
fibers. It is well established that thrombin concentrations have a major impact on fibrin structure, with increasing thrombin concentrations resulting in a denser fibrin network with thinner fibers [39–41]. Intriguingly, MASP-1 seems to have the opposite effect as increasing MASP-1 concentrations were associated with less dense fibrin networks and thicker fibers, suggesting that a mechanism other than increased thrombin generation may be involved. So far, however, the way MASP-1 alters clot characteristics remains unclear. Although we show that fibrin formation in plasma is mainly thrombin-dependent, MASP-1 may interfere with prothrombin and fibrinogen resulting in slower production of fibrin monomers. Lower polymerization rates are known to produce a looser network with thicker fibers [39]. It has also been suggested that FPB release from the fibrinogen β-chain promotes lateral aggregation resulting in formation of thick fibers [42,41]. Thus, additional FPB cleavage by MASP-1, as shown by Krarup et al. [19], may just have this effect. In addition, the observed differences in clot structure associated with MASP-1 may be related to different activation of FXIII and altered cross-linking of various proteins. These observations may also be in line with results by Gulla et al. [21] who suggested a similar clot composition but less cross-linking of clots made with MASP-1.

We show for the first time that MASP-1 is able to activate TAFI. This reaction is independent of thrombin generation as it was observed in both NCP and PT-DP and confirmed by SDS-PAGE. It should be noted, however, that MASP-1 activation of TAFI occurs at a much lower efficiency compared with thrombin/thrombomodulin. Nevertheless, additional TAFI activation by MASP-1 may explain the prolongation of lysis time associated with MASP-1 in the turbidimetric clotting and lysis assay, since TAFI is a major inhibitor of fibrinolysis. Another possible explanation for prolongation of lysis time may be the interaction between MASP-1 and FXIII which may lead to changes in fibrin cross-linking and cross-linking of antifibrinolytic proteins to fibrin, which in turn affects efficiency of fibrinolysis.

A less dense clot structure is generally associated with shorter lysis times, whereas we see longer lysis times of plasma clots made in the presence of MASP-1. Whether the profibrinolytic effect of a less dense network or the antifibrinolytic effect observed in the turbidimetric assay prevails in vivo remains to be investigated. If MASP-1 indeed led to prolongation of fibrinolysis, this would represent a further link between the complement system, in particular MBL pathway components, and changes in fibrin clot structure and prolongation of fibrinolysis, as our group has previously shown an independent association between complement C3 levels and prolongation of fibrinolysis [43–45].

A limitation of our study is the use of a recombinant mutant catalytic fragment of MASP-1 instead of the full-length molecule. We used this catalytic fragment for the following reasons: First of all, at present there is no appropriate method to purify native MASP-1 from plasma or serum and consequently several attempts to purify MASP-1 in sufficient amounts and free from contamination with other MASPs have failed. In addition, recombinant expression of full-length MASP-1 failed in all expression systems. In eukaryotic cells wild type MASP-1 is toxic. The recombinant MASP-1 constructs which could be expressed (e.g. in CHO cells) were mutants in the catalytic domain leading to reduced proteolytic efficiency. In contrast, the recombinant MASP-1 fragment we used in our study is the only preparation in the world which represents the wild type active human MASP-1. We are aware that, unlike the native protein, this catalytic fragment cannot form dimers or bind to MBL or ficolins and does not require an activation step and may therefore behave differently. However, the CUB-EGF-CUB domains, which are deleted in our fragment but present in the C1r, C1s and MASP proteins, are engaged in binding to the recognition molecules and consequently do not take part in substrate binding. The CCP modules, however, can contain exosites for substrate binding, as demonstrated for MASP-2 and C1s. Accordingly, our catalytic MASP-1 fragment contains both CCP modules. In addition, recombinant catalytic fragments of C1r, a MASp homolog, have been shown to retain the catalytic efficiency and substrate specificity of the entire molecule [9]. Endogenous MASP-1 circulates in plasma as a zymogen. The active recombinant fragment, rMASP-1 CCPI-CCP2-SP, we used in our study might activate the endogenous zymogen. In addition, it might also activate MASP-2 as recently shown [13]. However, in both cases the measured effects would still be due to MASP-1. Taken together, we do believe that the catalytic MASP-1 fragment we have used in our study represents the best model protein currently available to achieve our main goal which was to investigate MASP-1 catalytic properties towards coagulation proteins in plasma.

In summary, we used an array of coagulation factor activation and plasma clotting assays as well as SEM of plasma clots to show that MASP-1, albeit not able to induce fibrin clot formation completely independently of coagulation activation, interacts with plasma clot formation on different levels and influences the resulting fibrin structure. MASP-1 activates FXIII independently from thrombin and has a differential effect on FXIII Val34 and Leu34 variants. On the other hand, fibrinogen-to-fibrin conversion by MASP-1 is related to thrombin generation through prothrombin activation. Fibrin clot formation can be initiated by MASP-1 in plasma and this has an effect on lysis time, which may be related to changes in clot structure, activation of FXIII and TAFI, or other unknown factors. Our results support the first in vivo findings for a role of MASP-1 in coagulation and thrombus formation [22,23], but further investigations to elucidate the precise mechanisms and physiological relevance in humans are needed. Nevertheless, our results are consistent with a possible role for MASP-1 in plasma fibrin clot formation and fibrinolysis, which may have future clinical implications.

Acknowledgments

We would like to thank J.W. Weisel (University of Pennsylvania, Philadelphia, USA) for his advice on the interpretation of fibrin clot SEM images and S. Uitte de Willige (formerly at the University of Leeds, UK, now at the Erasmus University Medical Center, Rotterdam, The Netherlands) for valuable discussions.

Author Contributions

Conceived and designed the experiments: KH RA VS. Performed the experiments: KH RA FP VS. Analyzed the data: KH RA VS. Contributed reagents/materials/analysis tools: JD PG. Wrote the paper: KH RA VS. Revised the manuscript: KH RA JD PG VS.

References

1. Opal SM (2000) Phylogenetic and functional relationships between coagulation and the innate immune response. Crit Care Med 28: 877–80.

2. Markiewski MM, Nilsson B, Nilsson Eklblad K, Mullones TE, Lambris JD (2007) Complement and coagulation: strangers or partners in crime? Trends Immunol 28: 164–192.
5. Gál P, Barna L, Kocsis A, Za´vodszky P (2007) Serine proteases of the classical
complement pathway: similarities and differences. Immunobiology 212: 267–277.

8. Amara U, Flierl MA, Rittirsch D, Klos A, Chen Hui, et al. (2010) Molecular
biology of MBL-associated serine proteases (MASPs). Immunobiology 205: 467–475.

11. Møller-Kristensen M, Thiel S, Sjo¨holm A, Matsushita M, Jensenius JC (2007)
Targeting of mannan-binding lectin-associated serine protease-2 confers
protection from myocardial and gastrointestinal ischemia/reperfusion injury.
Proc Natl Acad Sci USA 104: 7529–7534.

14. Takahashi M, Ishida Y, Iwaki D, Kanno K, Suzuki T, et al. (2010) Essential role
of complement mannose-binding lectin-as sociated serine protease-1 in
and -2: a study on recombinant catalytic fragments. J Immunol 170: 1374–1382.

17. Presanis JS, Hajela K, Ambrus G, Ga´l P, Sim RB (2004) Differential substrate
specificity of human mannose-binding lectin-associated serine protease-1
and -2: a study on recombinant catalytic fragments. J Immunol 170: 1374–1382.

20. Gulla KC, Gupta K, Krarup A, Ga´l P, Schwaeble WJ, et al. (2010) Activation of
mannose-binding lectin-associated serine proteases leads to generation of a fibrin
clot. Immunology 129: 492–493.

23. La Bonte LR, Pavlov VI, Tan YS, Takahashi K, Takahashi M, et al. (2012)
Mannose-binding lectin-associated serine protease-1 is a significant contributor
to coagulation in a murine model of occlusive thrombosis. J Immunol 188:
485–491.

26. Collet JP, Allali Y, Leuty C, Tanay ML, Silvain J, et al. (2006) Altered fibrin
architecture is associated with hypofibrinolysis and premature coronary
afterthrombosis. Arterioscler Thromb Vasc Biol 26: 2567–2573.

29. Ajan R, Lim BCB, StandevenKF, Harrand D, Dolling S, et al. (2008) Common
variation in the C-terminal region of the fibrinogen β-chain affects fibrin
stability, fibrinolysis and clot rigidity. Blood 111: 643–650.

32. Krarup A, Mikkola H, Muszbek L, Laiho E, Syrjala¨M, Ha¨ma¨la¨inen E, et al. (1997)
Molecular mechanism of a mild phenotype in coagulation factor XIII (FXIII)
deficiency: a splicing mutation permitting partial correct splicing of FXIII A-
subunit mRNA. Blood 89: 1279–1287.

35. Jadhav MA, Lucas RC, Goldsberry WN, Maurer MC (2011) Design of factor
XIII V29F and V34L mutants of factor XIII (28–41) reveals roles of the P(9) and P(4) positions in
MSP-1 activation of the lectin complement pathway. J Immunol 185: 4169–4178.

38. Presanis JS, Hajela K, Ambrus G, Gál P, Sim RB (2004) Differential substrate
and inhibitor profiles for human MASP-1 and MASP-2. Mol Immunol 40:
921–929.

41. Dobi J, Harmat V, Beinrohr L, Bucsa I, Mogyeri M, et al. (2011) Cleavage of
kininogen and subsequent bradykinin release by the complement component:
linking activity. Thromb Haemost 80: 704.

44. Ryan EA, Mockros LF, Weisel JW, Lorand L (1999) Structural origins of fibrin
structure, fibrinolysis and clot rigidity. Blood 111: 643–650.

47. Hess K, Alzahrani S, Mathai M, Schroeder V, Carter AM, et al. A novel
mechanism for hypofibrinolysis in diabetes: The role of Component C3.
Diabetes. In press 2012.