Fibrinogen and clot-related phenotypes determined by fibrinogen polymorphisms: Independent and IL-6-interactive associations

H. Toine Cronje¹, Cornelie Nienaber-Rousseau¹, Lizelle Zandberg¹, Zelda de Lange¹, Fiona R. Green², Marlien Pieters¹*

¹ Centre of Excellence for Nutrition, North-West University, Potchefstroom, South Africa, ² Division of Cardiovascular Sciences, School of Medical Sciences, Faculty of Biology, Medicine & Health, University of Manchester, Manchester, United Kingdom

* marlien.pieters@nwu.ac.za

Abstract

Interleukin-6 (IL-6) induces the expression of fibrinogen, and polymorphic variation within the fibrinogen genes is believed to alter the magnitude of this expression. The identification of the functional relevance of individual fibrinogen single nucleotide polymorphisms (SNPs) has been hindered by the high linkage disequilibrium (LD) reported in the European fibrinogen gene locus. This study investigated two novel and 12 known fibrinogen SNPs of potential functional relevance, in 2010 Tswana individuals known to have low LD. We aimed to identify functional polymorphisms that contribute to clot-related phenotypes and total and γ' fibrinogen concentrations independently and through their interaction with IL-6, by taking advantage of the high fibrinogen and IL-6 concentrations and the low LD reported in black South Africans. Fibrinogen was significantly associated with IL-6, thereby mediating associations of IL-6 with clot formation and structure, although attenuating the association of IL-6 with clot lysis time. None of the common European fibrinogen haplotypes was present in this study population. Putative functional fibrinogen SNPs FGB–rs7439150, rs1800789 (−1420G/A) and rs1800787 (−148C/T) were significantly associated with fibrinogen concentration and altered clot properties, with several associations significantly influenced by IL-6 concentrations. The impact of harbouring several minor fibrinogen SNP alleles on the association of IL-6 and fibrinogen concentration was cumulative, with possession of each additional minor allele showing a stronger relationship of IL-6 with fibrinogen. This was also reflected in differences in clot properties, suggesting potential clinical relevance. Therefore, when investigating the effect of fibrinogen genetics on fibrinogen concentrations and CVD outcome, the possible interactions with modulating factors and the fact that SNP effects seem to be additive should be taken into account.
Introduction

Fibrinogen is central to blood coagulation and the inflammatory response. As a haemostatic protein, fibrinogen is activated by thrombin to form fibrin, the main constituent of a blood clot [1]. As an acute phase reactant, fibrinogen is up-regulated upon stimulation by inflammatory cytokines, primarily interleukin-6 (IL-6), in response to physiological trauma such as infection/inflammation [2]. Both fibrinogen and IL-6 are prospectively associated with cardiovascular disease (CVD) risk [3–5], although causality is under debate [6–9].

The fibrinogen phenotype is heritable and several single nucleotide polymorphisms (SNPs) within the fibrinogen α, β and γ chain genes (FGA, FGB, FGG) have been identified as contributors to this heritability in Europeans [10–13]. Fibrinogen expression is regulated on two levels: under basal conditions and during the acute phase response [14, 15]. This acute phase-induced increase in fibrinogen is largely mediated by IL-6, through the JAK/STAT pathway. In addition, sequences responsive to IL-6, and crucial for full IL-6-induced fibrinogen expression, have been identified upstream of the fibrinogen genes [14, 15].

Genetic variation in the fibrinogen genes is hypothesised to alter the magnitude of fibrinogen expression in response to IL-6 [13, 16, 17]. Thus far, the focus of investigations has been on polymorphisms within the FGB promoter region that have been implicated in both epidemiological [16] and in vitro [17] studies to interact with IL-6 in influencing fibrinogen concentrations. Greater IL-6-induced fibrinogen expression has been reported for the FGB rs1800790A (–455A) and rs1800791A (–854A) [16–20], while the opposite has been observed for FGB rs1800789 (–1420G) and rs1800787 (–148T) [16, 17]. Apart from the FGB polymorphisms, IL-6 interacted with FGA rs2070011T in vivo to enhance fibrinogen expression, whereas FGA rs6050, FGG rs2066865 and FGG rs1049636 had no effect [13].

In addition to regulating total fibrinogen concentration, IL-6 is able to up-regulate the production of γ' fibrinogen, a common splice variant, comprising 8 to 15% of total fibrinogen concentration [21–23]. IL-6 specific responsive elements have been characterised in the FGG promoter region [24, 25], and in vitro research reported greater increases in γ' compared to total fibrinogen in the presence of IL-6, suggesting alterations to the alternative splicing of the γ fibrinogen gene during inflammation [22]. Elevated fibrinogen γ' concentration is a risk factor for arterial thrombosis, although research on the individual and IL-6-interactive effects of fibrinogen polymorphisms predicting γ' fibrinogen concentrations is limited [26–29]. Increased IL-6, total and γ’ fibrinogen are also independently associated with altered clot properties, including faster clot formation, increased fibrin density, thinner fibrin fibres and decreased clot permeability [30–34]. The functional effects of the interactions between genetic variants and IL-6 on clot properties have not been described previously.

Tight LD within the fibrinogen SNPs in Europeans has made the identification of functional SNPs difficult. Studies have implicated the FGB -455G/A, -854G/A and -148C/T polymorphisms to be functional in Europeans, although findings from epidemiological [16] and experimental work [17] have been contradictory. The African population is known for its complex LD pattern and great genetic diversity, more so than any other population [35, 36]. Furthermore, increased fibrinogen and IL-6 concentrations have been observed in black South Africans when compared to their white counterparts [37, 38]. The South African arm of the prospective urban and rural epidemiology (PURE) study indicated low LD together with markers indicative of chronic low-grade inflammation—as reported by increased fibrinogen, IL-6, homocysteine and C-reactive protein [39, 40]—making this cohort ideal for the investigation of independent and IL-6-regulated gene interactions. Therefore, this study investigates selected fibrinogen polymorphisms in terms of their independent and IL-6-interactive associations with total and γ' fibrinogen concentrations, and the downstream functional effects in
terms of clot formation, structure and lysis utilizing the unique Tswana population. Our results contribute to the broader understanding of potential factors that regulate fibrinogen-related, clinically relevant phenotypes.

Materials and methods

Study population and ethical considerations

This study is affiliated with the international PURE [41] study and is a cross-sectional investigation of the baseline data collected in South Africa. Details pertaining to participant selection and recruitment were published previously [39]. In short, the study population consisted of 2010 apparently healthy, self-identified black Tswana-speaking South African adults between the ages of 35 and 70 years. Participants provided voluntary written informed consent prior to participating in the study, in addition to specific consent for genetic analysis. Anonymity of participants remains ensured by providing only coded electronic data containing no information that can be used to identify individuals. Ethical approval for the research was obtained from the Health Research Ethics Committee of the North-West University (04M10 and NWU-00058-16-S1) and the study was conducted in accordance with the revised version (2000) of the Helsinki Declaration of 1975.

Blood collection and storage

Fasting blood samples were collected between 7:00 and 11:00 am, and centrifuged within 30 minutes of collection at 2000 x g for 15 minutes. Blood samples for haemostatic variables were collected in 3.2% sodium citrate tubes, for lipid and IL-6 analyses in tubes without anti-coagulants and for glycated haemoglobin (HbA1c) in sodium fluoride tubes. Following centrifugation, samples were snap-frozen and stored at –80˚C until analysis.

Biochemical analyses

HbA1c concentrations were quantified using a hexokinase method from Synchron® systems (Beckman Coulter Co., Fullerton, CA, USA), and the D-10 haemoglobin testing system (Bio-Rad, Hercules, California, USA). A Sequential Multiple Analyser Computer, using the Konelab™ auto-analyser (Konelab 20i, Thermo Fisher Scientific, Vantaa, Finland) was used to measure high-density lipoprotein cholesterol (HDL-c) concentrations. Plasminogen activator inhibitor type-1 activity (PAI-1 act) was quantified by an indirect enzymatic method (Spectrolyse PAI-1, Trinity Biotech, Bray, Ireland). Serum IL-6 was measured by means of the Elecsys, through ultra-sensitive enzyme immunoassays (Elecsys 2010, Roche, Basel, Switzerland).

Total fibrinogen concentrations were quantified using the modified Clauss method on the Dade Behring BCS coagulation analyser (Multifibrin U-test Dade Behring, Deerfield, IL, USA). An enzyme-linked immunosorbent assay using a 2G2.H9 mouse monoclonal coating antibody against human γ’ fibrinogen (Santa Cruz Biotechnology, Santa Cruz, USA) and a goat polyclonal horseradish peroxidase-conjugated antibody against human fibrinogen (Abcam Cambridge, USA) was used to measure γ’ fibrinogen concentrations. Fibrinogen γ’ is expressed both as absolute concentration and as a percentage of total fibrinogen (%γ’).

Plasma fibrinolytic potential was determined by measuring turbidity with a spectrophotometer (A405) [42]. Tissue plasminogen activator (tPA; Actilyse, Boehringer Ingelheim, Ingelheim, Germany) was added to plasma clots induced by tissue factor (TF; Dade Innovin, Siemens Healthcare Diagnostics Inc., Marburg, Germany) according to the method of Lisman et al. [43]. The tPA and TF concentrations were modified to obtain a clot lysis time (CLT) between 60 and 100 minutes in the external control sample (pooled plasma) included in each
run. Final concentrations in the plasma clots were 125 x diluted TF (approximately 59 pmol/L), 100 ng/mL tPA, 17 mmol/L CaCl₂, and 10 mmol/L phospholipid vesicles (Rossix, Mölnadal, Sweden). Resultant turbidity curves were analysed using Origin® software version 8.5 (Origin lab®, 2010). CLT (minutes) was calculated as the difference between the time at the midpoint of clear and maximum turbidity (clot formation) and the midpoint between maximum and clear turbidity (clot lysis). In addition, lag time (minutes) was calculated as an indicator of the time required for the activation of the coagulation cascade by TF and for protofibrils to reach sufficient length to allow lateral aggregation. The slope of the curve (x10⁻³ au/s) during clot formation was used as a representation of the rate of lateral aggregation of the fibrin protofibrils. Lastly, maximum absorbance, the increase in absorbance at the peak of the curve (nm), was used as an indicator of fibre diameter.

DNA isolation, SNP selection and genotyping

Genomic DNA was extracted from buffy coat using the QIAGEN FlexiGene™ kit. The quality of the DNA was determined using the NanoDrop™ spectrophotometer (ND-1000, Wilmington, DE, USA). Fourteen SNPs, spanning the three fibrinogen genes, were selected for genotyping based on literature identifying them as the top SNPs to demonstrate independent and IL-6-induced associations with total or γ' fibrinogen concentrations [9, 11, 13, 44–49]. Furthermore, the promoter region of FGB was sequenced in a subgroup of 28 randomly selected individuals for the identification of novel SNPs. ABI Prism®, BIGDye® Terminator version 3.1 Ready Reaction Cycle Sequencing Kits (Applied Biosystems, CityFoster, CA, USA) were used, and electropherograms were aligned using BioEdit (version 7.1.3.0, Ibis Biosciences, Carlsbad, CA, USA).

The polymorphisms selected for genotyping were FGB-rs7439150, rs2227385 (novel), rs1800789 (-1420G/A), rs2227388 (novel), rs1800791 (-854G/A), rs1800790 (-455G/A), rs1800788 (-249C/T), rs1800787 (-148C/T), rs4220, rs4463047, FGA-rs6050, rs2070011 (2224G/A), and FGG-rs2066865 and rs1049636 (9340T/C). Three methods were used to genotype these polymorphisms, of which two (Thermo Fischer Scientific®, Taqman based assays and the Illumina® VeraCode GoldenGate assay technology using a BeadXpress® platform), have been described previously [39]. In addition, FGB-rs1800789, rs1800790, rs4463047 and rs7439150 were genotyped by competitive allele-specific polymerase chain reactions (KASP) with supplies obtained from LGC Limited. Custom-designed assays and synthetic controls were manufactured by KBioscience and are presented in S1 Table (LGC, Middlesex, TW11 0LY, UK).

A two-step 61–to–55°C touchdown polymerase chain reaction (PCR) protocol was performed in a Hydrocycler 4™ water bath thermal cycler (LGC, Middlesex, TW11 0LY, UK). The fluorescent signal of the PCR products was measured by a FLUOstar Omega SNP plate reader (BMG LABTECH Ltd) and the data were analysed by means of KlusterCaller™ V. 3.4.1.36 software (LGC, Middlesex, TW11 0LY, UK).

Statistical analyses

The statistical analyses were performed in three phases. Firstly, the SNPs were investigated in terms of their location, LD and haplotypes using the Ensembl database release 84 [50] and Haploview version 4.2 [51]. In addition, their independent associations with phenotype outcomes (total and γ fibrinogen concentration, lag time, slope, maximum absorbance and CLT) were determined by independent t-tests and analysis of co-variance (ANCOVA) adjusting for age, gender, body mass index (BMI), human immunodeficiency virus (HIV) status, HbA1c and HDL-c. When CLT was used as an outcome variable, it was also adjusted for PAI-1 act.

The second phase involved investigating the association of the fibrinogen phenotypes with IL-6 independent from the polymorphisms. Differences in phenotypes were tested with
analysis of variance (ANOVA) and ANCOVA using IL-6 stratified by quartiles as the categorical variable. Tukey’s honest significant difference post-hoc tests were performed to determine significant inter-group differences.

Lastly, interaction effects between IL-6 and the fibrinogen polymorphisms on the fibrinogen phenotypes were determined by creating interaction terms and entering them into an ANCOVA with full factorial analysis. Adjustments were made for the same covariates listed above. Multiple testing was accounted for by performing Benjamini and Hochberg adjustments. Q-Q plots were used to evaluate the normality of the standardised residuals. Interaction data are reported as the slope of the linear regression line between IL-6 and the fibrinogen phenotypes split according to genotype.

In order to determine the possibility of a cumulative effect of the SNPs on the IL6-fibrinogen relationship, polymorphisms indicating significant interactions were grouped by means of the generation of a simple genetic ‘risk score’. Each genotype forming part of the model (significant interaction upon adjustment for multiple testing), was given a score per individual genotype. A score of zero was given for major allele homozygotes; one was given to the heterozygotes and two to minor allele homozygotes. Where the minor allele frequency (MAF) was low and only two groups were compared (see below), a score of one was given to all minor allele carriers.

Based on the minor allele frequency (determined by 1 out of 28 individuals showing variation at any given SNP in the FGB promoter region, upon sequencing) the power calculation indicated that in order to obtain 80% power to detect a medium effect size (Cohen’s d-value of 0.3), the genotype groups should contain a minimum of 71 individuals [52]. Analyses for SNPs having fewer than 71 individuals in the minor allele homozygote group were, therefore, performed using two groups only, combining heterozygotes and minor allele homozygotes.

Any analyses (t-tests, ANOVA, full factorial analysis ANCOVA) yielding significant results in terms of total or γ fibrinogen were followed by adjustments for these proteins during statistical testing using clot properties as outcome variables. These adjustments were made to identify SNP/IL-6-clot property outcomes that were not mediated by total and/or γ fibrinogen concentration. The statistical package for the social sciences (SPSS®) version 23 (IBM® Corp, 2015) and Statistica® version 13 (Statsoft Inc., Tulsa, OK, USA) were used for the above described analyses.

**Results**

**Description of individual polymorphisms**

Ten of the 14 investigated polymorphisms are situated in and around the β chain gene, of which eight are within the promoter area, one in exon 8 (Arg478Lys, previously reported as Arg448Lys) and one in the 3’ untranslated region (UTR). Two α chain variants, one in exon 2 (Thr331Ala, previously reported as Thr312Ala) and one in the promoter area, and two γ chain variants, both in the 3’ UTR, are also included. The relative positions of these SNPs, spanning a 50 Kb region on chromosome 4, are illustrated in Fig 1. All the variants were in Hardy-Weinberg equilibrium.

**LD and haplotype construction**

Fig 2 depicts the LD pattern (illustrated by D’ values upon an r^2 colour scheme) observed for the 14 SNPs. Lower recombination rates are indicated by increased numerical values (D’) and darker coloured blocks (r^2). Blocks containing no numerical value indicate a D’ of 1.0. Two haplotype blocks FGB-rs7439150’rs2227385’rs1800789 and FGG rs2066865’rs1049636 were constructed via the method suggested by Gabriel et al. [53]. No complete LD (D’ and r^2 = 1.0) was observed, as the LD pattern was disturbed owing to differing MAFs that led to relatively
low $r^2$ values. Acknowledging the limited LD, all further analyses were performed using individual polymorphisms only.

**Associations of individual polymorphisms with fibrinogen-related phenotypes**

Population characteristics and associations of ten of the 14 SNPs with fibrinogen-related phenotypes have been reported previously [39] and are shown in the online supporting

![Image](https://doi.org/10.1371/journal.pone.0187712.g001)

**Fig 1. Fourteen polymorphisms spanning the fibrinogen gene cluster.** Image generated through the Ensembl database [50].

![Image](https://doi.org/10.1371/journal.pone.0187712.g002)

**Fig 2. Pairwise LD structure of 14 fibrinogen SNPs, illustrated by $D'$ values on an $r^2$ colour scheme.** Empty boxes indicate $D'$ values of 1.0. Increased numerical values indicate stronger evidence of LD. Image generated through Haploview software [51].
information only (S2, S3 and S4 Tables). Briefly, $FGB$–854$A$ and $FGG$–rs1049636$C$ were positively associated with total fibrinogen ($p = 0.04$ and $0.0009$), and $FGA$–rs2070011$A$ with higher $\gamma'$ fibrinogen concentrations ($p = 0.008$). In terms of clot properties, $FGB$–148$T$ was associated with a larger fibre diameter ($p = 0.001$), whereas $FGA$–rs2070011$A$ and $FGG$–rs1049636$C$, lost significance in terms of their effect on maximum absorbance (fibre diameter) upon adjustment for $\gamma'$ and total fibrinogen, respectively ($p = 0.06$ and $0.33$).

Associations of each of the four newly investigated promoter region SNPs with the outcome phenotypes are shown in Table 1. $FGB$–1420$A$ was significantly associated with total fibrinogen concentrations ($p = 0.02$) and the same trend was observed for $FGB$–rs7439150$A$ and $FGB$–455$A$ ($p \approx 0.05$). Larger fibre diameter and longer CLT were observed in the presence of the $FGB$–1420$A$ allele ($p = 0.04$ and $0.02$). In addition, $FGB$–455$A$ and $FGB$–rs7439150$A$ were positively associated with $\gamma'$ fibrinogen concentrations ($p = 0.02$ and $0.005$), and $FGB$–rs7439150$A$ with fibre diameter ($p = 0.02$). Associations with clot properties were largely mediated by the primary fibrinogen associations, as most of the significance was lost upon adjustment for total and/or $\gamma'$ fibrinogen, apart from the association between $FGB$–1420$A$ and CLT, which remained ($p = 0.03$).

The association of IL-6 with fibrinogen-related phenotypes

The basic descriptive characteristics of the study population have been reported in a previous publication [39]. Descriptive statistics for fibrinogen and clot properties, including their

<table>
<thead>
<tr>
<th>SNP pseudonym</th>
<th>Genotype</th>
<th>$FGB$–rs7439150$A$</th>
<th>$FGB$–1420 $G&gt;A$</th>
<th>$FGB$–455 $G&gt;A$</th>
<th>$FGB$–rs4463047</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minor allele frequency (%)</td>
<td>major allele</td>
<td>$A = 6.95$</td>
<td>$A = 6.78$</td>
<td>$A = 3.32$</td>
<td>$T = 11.4$</td>
</tr>
<tr>
<td>Genotype group size (n)</td>
<td>minor allele carrier</td>
<td>$GG^1 (1548)$</td>
<td>$GG^2 (1513)$</td>
<td>$GG^1 (1652)$</td>
<td>$CC^1 (1424)$</td>
</tr>
<tr>
<td>Total fibrinogen (g/L)</td>
<td>major allele</td>
<td>$3.66 \pm 1.23$</td>
<td>$3.64 \pm 2.12$</td>
<td>$3.66 \pm 2.12$</td>
<td>$3.71 \pm 2.18$</td>
</tr>
<tr>
<td>Fibrinogen $\gamma'$ (mg/mL)</td>
<td>minor allele</td>
<td>$3.97 \pm 2.29$</td>
<td>$3.96 \pm 2.29$</td>
<td>$4.05 \pm 2.34$</td>
<td>$3.52 \pm 1.99$</td>
</tr>
<tr>
<td>Fibrinogen $\gamma'$ (%)</td>
<td>major allele</td>
<td>$0.37 \pm 0.25$</td>
<td>$0.38 \pm 0.25$</td>
<td>$0.38 \pm 0.26$</td>
<td>$0.38 \pm 0.27$</td>
</tr>
<tr>
<td>Lag time (min)</td>
<td>minor allele</td>
<td>$12.1 \pm 8.07$</td>
<td>$12.3 \pm 8.27$</td>
<td>$12.1 \pm 8.01$</td>
<td>$12.1 \pm 8.27$</td>
</tr>
<tr>
<td>Slope ($x10^{-3}$ au/s)</td>
<td>minor allele</td>
<td>$6.67 \pm 1.97$</td>
<td>$6.47 \pm 1.96$</td>
<td>$6.48 \pm 1.99$</td>
<td>$6.47 \pm 1.97$</td>
</tr>
<tr>
<td>Maximum absorbance (nm)</td>
<td>major allele</td>
<td>$9.58 \pm 4.42$</td>
<td>$9.54 \pm 4.35$</td>
<td>$9.65 \pm 4.36$</td>
<td>$9.67 \pm 4.41$</td>
</tr>
<tr>
<td>Clot lysis time (min)</td>
<td>minor allele</td>
<td>$10.0 \pm 3.99$</td>
<td>$10.3 \pm 4.09$</td>
<td>$9.75 \pm 3.95$</td>
<td>$9.60 \pm 4.32$</td>
</tr>
</tbody>
</table>

$A$ = adenine; $C$ = cytosine; $FGB$ = fibrinogen beta chain gene; $G$ = guanine; $rs$ = reference sequence; $T$ = thymine; Lag time = time required for the activation of the coagulation cascade by TF and for protofibrils to reach sufficient length to allow lateral aggregation; Slope = rate of lateral aggregation of fibrin protofibrils; Maximum absorbance = indicator of fibre diameter.

Data presented as mean $\pm$ SD.

*p* $< 0.05$

* $p$ $< 0.05$ upon adjustment for total and $\gamma'$ fibrinogen

* Strong evidence of LD ($r^2 = 0.89$; $D'$ = 0.95).

Age, gender, BMI, HIV-status, HbA1c, and HDL-c were covariates. Associations with CLT were adjusted for PA-I1 act additionally.

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Table 2. Outcome phenotypes descriptive statistics and association with IL-6- quartiles.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Whole group (n)</th>
<th>Interleukin-6 quartiles (pg/mL)</th>
<th>p-value unadjusted (adjusted)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Quartile 1 &gt; 0.76</td>
<td>Quartile 2 1.50–2.84</td>
</tr>
<tr>
<td>Total fibrinogen (g/L)</td>
<td>3.69 ± 2.18 (1593)</td>
<td>3.02 ± 1.62</td>
<td>3.32 ± 1.86</td>
</tr>
<tr>
<td>Fibrinogen γ' (mg/L)</td>
<td>0.38 ± 0.27 (1591)</td>
<td>0.32 ± 0.22</td>
<td>0.36 ± 0.25</td>
</tr>
<tr>
<td>Fibrinogen γ' (%)</td>
<td>12.1 ± 8.25 (1527)</td>
<td>12.5 ± 8.56</td>
<td>12.4 ± 7.58</td>
</tr>
<tr>
<td>Lag time (min)</td>
<td>6.46 ± 1.97 (1480)</td>
<td>6.24 ± 1.99</td>
<td>6.46 ± 2.03</td>
</tr>
<tr>
<td>Slope (x10⁻³ au/s)</td>
<td>9.70 ± 4.42 (1492)</td>
<td>9.19 ± 3.80</td>
<td>9.60 ± 4.37</td>
</tr>
<tr>
<td>Maximum absorbance (nm)</td>
<td>0.43 ± 0.16 (1445)</td>
<td>0.41 ± 0.14</td>
<td>0.43 ± 0.15</td>
</tr>
<tr>
<td>CLT (min)</td>
<td>57.3 ± 11.2 (1591)</td>
<td>58.3 ± 10.8</td>
<td>56.9 ± 10.9</td>
</tr>
</tbody>
</table>

Data presented as mean ± SD

Adjusted p-value = adjusted for total and γ' fibrinogen; CLT = clot lysis time; Lag time = time required for the activation of the coagulation cascade by TF and for protofibrils to reach sufficient length to allow lateral aggregation; Slope = rate of lateral aggregation of fibrin protofibrils; Maximum absorbance = indicator of fibre diameter.

Means with the same symbol differ significantly for individual outcome variables

Means with the same numerical value differ significantly for individual outcome variables upon adjustment.

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association with IL-6 (presented as quartiles), are shown in Table 2. The mean IL-6 concentration was 6.50 ± 21.0 pg/mL. Total and γ' fibrinogen concentrations were positively, although not linearly, associated with IL-6 with significantly higher concentrations in the fourth IL-6 quartile compared to the first three quartiles. The % γ' fibrinogen did not reach significance, implying that the fibrinogen γ' association is probably a reflection of the association of IL-6 with total fibrinogen. Positive associations were also observed for lag time, slope and maximum absorbance, although subsequent adjustment for total fibrinogen concentration attenuating this effect. Adjustments for total and γ' fibrinogen, and thereafter PAI-1act (as a main modulator of CLT) significantly increased this negative association (Fig 3).

Genotype-IL-6 interactions in terms of phenotype predictions

Interaction analyses were performed for each polymorphism with IL-6 in relation to each of the fibrinogen measures and clot properties. The association between IL-6 and fibrinogen phenotype, stratified according to genotype for each of the significant polymorphisms is reported in Tables 3 and 4. IL-6 concentration was associated with a steeper increase in fibrinogen concentration in minor allele carriers of seven SNPs and a shallower increase in total fibrinogen in FGA-rs6050. For FGG-rs2066865, a shallower increase was observed in the presence of one minor allele, with a steeper increase in the presence of two minor alleles, more so than for the common allele homozygotes. Upon adjusting for multiple testing, IL-6 did not significantly interact with any of the investigated SNPs to influence fibrinogen γ'. IL-6 interacted with six SNPs in determining fibre diameter (maximum absorbance), in which the presence of the minor allele resulted in a stronger positive association between IL-6 and maximum absorbance than the respective major alleles. Adjustment for fibrinogen concentration largely nullified
these associations, with only two associations remaining significant (FGB-rs7439150 and –148C/T). FGB–1420G/A, which is in high LD with FGB-rs7439150 and –148C/T, however, lost its significance after adjustment for fibrinogen owing to its independent association with fibrinogen concentration. No significant interactions were observed for lag time, slope or CLT. Removal of individuals with IL-6 concentrations higher than 100 pg/ml (n = 8) resulted in a loss of significance for many of the interactions presented in Tables 3 and 4. The only remaining IL-6 interaction was with FGG-rs1049636 in terms of fibrinogen concentration [p < 0.001, slope 0.056 (0.041–0.071) for the major allele carriers and slope 0.110 (0.082–0.138) for the minor allele carriers].

In addition to the individual interaction analyses, the seven variants resulting in a steeper fibrinogen increase in the presence of IL-6 (Table 3) were grouped in a genetic ‘risk score’ to determine whether there were additive effects when these genotypes occurred together in an individual. Each individual’s score was composed of the sum of the seven polymorphisms allocated a value of either zero (major allele homozygote) or one (minor allele carrier). Final scores ranked from zero to six, therefore, scores represented the presence of zero to six minor allele groups. The risk score showed a significant interaction with IL-6 (p < 0.001) in the prediction of fibrinogen concentrations. Fig 4 schematically represents the slope the association between IL-6 and fibrinogen for each score. The addition of a minor allele group to the risk score increased the slope of the IL-6–fibrinogen association. The fifth and sixth risk score were removed from Fig 4 as the sizes of these groups were too small to provide adequate power (n = 24 and 5, respectively). The interaction, however, remained significant upon removal of these groups (p < 0.001). The regression coefficient for the line indicating the IL-6–fibrinogen association for risk score one differed significantly from risk score zero (p = 0.04), with two, three and four differing from one (p = 0.01, 0.048, 0.005, respectively) but not from each other, indicating a possible threshold to the additive effect when more than three risk alleles occurred together.

**Discussion**

High LD in the fibrinogen gene cluster in Europeans continues to hinder the identification of functional polymorphisms contributing to fibrinogen concentration and functionality...
This becomes even more complex when taking the role of transcriptional enhancers, such as IL-6, into consideration. Although little is known about the fibrinogen genes in Africans, preliminary evidence suggests distinct genetic differences from that of Europeans, including higher recombination rates and polymorphic variance. In

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Gene</th>
<th>IL-6 interaction with SNP</th>
<th>Interaction p-value</th>
<th>Major allele homozygotes</th>
<th>Heterozygotes or minor allele carriers (where only two groups)</th>
<th>Minor allele homozygotes (where three groups)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total fibrinogen (g/L)</td>
<td>FGB</td>
<td>rs7439150&lt;sup&gt;5&lt;/sup&gt;</td>
<td>0.001</td>
<td>GG 1293</td>
<td>0.015 (0.009–0.020)</td>
<td>174</td>
</tr>
<tr>
<td></td>
<td>FGB</td>
<td>rs1800789&lt;sup&gt;5&lt;/sup&gt;</td>
<td>&lt;0.001</td>
<td>GG 1263</td>
<td>0.012 (0.006–0.018)</td>
<td>240</td>
</tr>
<tr>
<td></td>
<td>FGB</td>
<td>rs1800788</td>
<td>&lt;0.001</td>
<td>CC 1419</td>
<td>0.017 (0.012–0.023)</td>
<td>154</td>
</tr>
<tr>
<td></td>
<td>FGB</td>
<td>rs1800787&lt;sup&gt;5&lt;/sup&gt;</td>
<td>0.001</td>
<td>CC 1164</td>
<td>0.016 (0.010–0.022)</td>
<td>152</td>
</tr>
<tr>
<td></td>
<td>FGB</td>
<td>rs4220</td>
<td>0.003</td>
<td>GG 1121</td>
<td>0.016 (0.010–0.022)</td>
<td>224</td>
</tr>
<tr>
<td></td>
<td>FGA</td>
<td>rs6050</td>
<td>&lt;0.001</td>
<td>AA 676</td>
<td>0.039 (0.028–0.049)</td>
<td>528</td>
</tr>
<tr>
<td></td>
<td>FGA</td>
<td>rs2070011</td>
<td>0.010</td>
<td>GG 925</td>
<td>0.015 (0.009–0.021)</td>
<td>416</td>
</tr>
<tr>
<td></td>
<td>FGG</td>
<td>rs2066865</td>
<td>&lt;0.001</td>
<td>CC 746</td>
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<tr>
<td></td>
<td>FGG</td>
<td>rs1049636</td>
<td>&lt;0.001</td>
<td>TT 975</td>
<td>0.014 (0.008–0.020)</td>
<td>371</td>
</tr>
</tbody>
</table>

A = adenine; C = cytosine; Cl = confidence interval; G = guanine; IL-6 = interleukin-6; rs = reference sequence; T = thymine; rs1800789 = -1420 G/A; rs1800788 = -249 C/T; rs1800787 = -148 C/T; rs1800790 = -555 G/A; rs2070011 = 2224 G/A; rs1049636 = 9340 T/C

<sup>5</sup>Variants in high linkage disequilibrium (r<sup>2</sup> > 0.82; D' > 0.92)

<sup>*</sup>In the regression line y = mx + c, “slope” refers to m.

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Fibrinogen-related phenotypes determined by independent and IL-6-interactive fibrinogen SNP associations

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Gene</th>
<th>IL-6 interaction with SNP</th>
<th>Interaction p-value</th>
<th>Major allele homozygotes</th>
<th>Heterozygotes or minor allele carriers (where only two groups)</th>
<th>Minor allele homozygotes (where three groups)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum absorbance (nm)</td>
<td>FGB</td>
<td>rs7439150&lt;sup&gt;6&lt;/sup&gt;</td>
<td>&lt;0.001</td>
<td>0.01</td>
<td>GG 1274</td>
<td>1x10&lt;sup&gt;-4&lt;/sup&gt;(-3x10&lt;sup&gt;-4&lt;/sup&gt;–0.001)</td>
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<tr>
<td></td>
<td>FGB</td>
<td>rs1800789&lt;sup&gt;6&lt;/sup&gt;</td>
<td>&lt;0.001</td>
<td>0.82</td>
<td>GG 1245</td>
<td>2x10&lt;sup&gt;-4&lt;/sup&gt;(-3x10&lt;sup&gt;-4&lt;/sup&gt;–0.001)</td>
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<tr>
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<td>FGB</td>
<td>rs1800788</td>
<td>0.001</td>
<td>0.04</td>
<td>CC 1407</td>
<td>0.001 (2x10&lt;sup&gt;-4&lt;/sup&gt;–0.001)</td>
</tr>
<tr>
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<td>rs1800787&lt;sup&gt;6&lt;/sup&gt;</td>
<td>&lt;0.001</td>
<td>0.01</td>
<td>CC 1157</td>
<td>1x10&lt;sup&gt;-4&lt;/sup&gt;(-3x10&lt;sup&gt;-4&lt;/sup&gt;–0.001)</td>
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<td>FGA</td>
<td>rs6050</td>
<td>0.001</td>
<td>0.04</td>
<td>AA 663</td>
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<td>0.003</td>
<td>0.23</td>
<td>TT 970</td>
<td>0.001 (3x10&lt;sup&gt;-4&lt;/sup&gt;–0.001)</td>
</tr>
</tbody>
</table>

A = adenine; C = cytosine; Cl = confidence interval; G = guanine; IL-6 = interleukin-6; rs = reference sequence; T = thymine; rs1800789 = -1420 G/A; rs1800788 = -249 C/T; rs1800787 = -148 C/T; rs1800790 = -555 G/A; rs2070011 = 2224 G/A; rs1049636 = 9340 T/C

Adjusted = adjusted for total fibrinogen (g/L); Slope = rate of lateral aggregation of fibrin protofibrils; Maximum absorbance = indicator of fibre diameter.

<sup>6</sup>Variants in high linkage disequilibrium (r<sup>2</sup> > 0.82; D' > 0.92)

<sup>*</sup>In the regression line y = mx + c, “slope” refers to m.

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addition, this population is known for altered fibrinogen and IL-6 phenotypes. This study, therefore, aimed to identify possible functional fibrinogen SNPs and novel IL-6-interactions related to total and γ fibrinogen, as well as indicators of clot formation, structure and lysis by using the unique African genetic and phenotypic profile. Our data revealed that none of the common European fibrinogen haplotypes is present in this African population. We demonstrate, in a population with chronic low-grade inflammation, that IL-6 interacted with several of the fibrinogen SNPs to influence fibrinogen concentration and even resulted in altered fibrin clot properties. A novel finding was that IL-6 not only interacted with individual SNPs, but that an additive effect was also observed when harbouring more than one risk allele concurrently.

The MAF of 12 of the 14 investigated SNPs was significantly lower in this Tswana population than that reported globally [50]. No variation has been reported for the FGB- rs2227385 and FGB-rs2227388 SNPs outside Africa [50]. As predicted, a high recombination rate was observed in this study population, resulting in no complete LD among any of the SNPs. However, strong LD between FGB-1420G/A and -148C/T was observed, similar to the finding of Dehghan et al. [54], although the LD observed for FGB-rs7439150 with these SNPs is novel. This lack of full LD differs from numerous reports of almost complete LD between the –1420G/A, –933C/T, –455G/A and –148C/T SNPs in Europeans [11, 16, 17, 49, 55–57]. None of the frequently occurring haplotypes, consisting of –1420G/A, –933C/T, –854G/A, –455G/A, –249G/A and –148C/T, [11, 16, 17], was present in this study population, highlighting the genetic diversity in Africans. In addition, FGB-933C/T revealed no genetic variation in this population.

Fig 4. Association of total fibrinogen concentrations with circulating interleukin-6 by risk score groups.

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In the present study, FGB –854A, –1420A and FGG-rs1049636C were associated with higher total fibrinogen concentrations. Higher γ’ fibrinogen concentrations (but not % γ’) were observed in the presence of FGB –455A, rs7439150A and FGA-rs2070011A. Furthermore, a positive association between FGB –148T and fibre diameter, and FGB –1420A and CLT was observed. IL-6 correlated positively with total and γ’ fibrinogen, thereby accelerating clot formation and increasing fibre diameter. CLT was, however, negatively associated with IL-6, and both total and γ’ fibrinogen concentration attenuated this association. Five FGB, one FGA and one FGG SNP significantly interacted with circulating IL-6 to steepen the IL-6 fibrinogen association. These interactions proved to be additive, with the presence of more than one minor allele across the gene cluster resulting in greater increases in IL-6-induced fibrinogen expression, although a threshold was reached when more than three minor allele variants occurred together in an individual. Lastly, these interactive associations reflected functional effects in terms of the rate of lateral aggregation and fibre diameter, thus indicating a possible mechanism by which the fibrinogen SNPs, during the acute phase, could enhance thrombotic risk.

Individual SNP associations for ten of the polymorphisms investigated have been reported previously [39]. The associations of the four additionally genotyped SNPs, although not all reaching significance, are in agreement with the existing literature. Increased fibrinogen concentrations in the presence of FGB-rs7439150A, –1420A and –455A have been reported in case-control and genome-wide association analyses [9, 54, 56, 58–65]. FGB-rs4463047T was negatively associated with total fibrinogen concentrations in three association studies [9, 61, 66], whereas we did not observe any association at this locus. The novel association of –1420A with faster CLT, independent of fibrinogen concentration, deserves further investigation.

Most of the research into both the heritability of and variant associations with fibrinogen has been conducted in European populations. Although limited information is available for African-Americans [9, 66, 67], being an ad-mixed population, their genetic background also differs from individuals from continental Africans. The lack of LD and common haplotypes, alongside the inability to reproduce results of independent SNP associations in this current investigation, questions whether heritability estimates obtained in Europeans or African Americans can be extrapolated to Africans. Future studies should replicate the existing heritability studies in African ethnic sub-groups to address this. The 14 SNPs investigated here contributed a mere 0.5% to the variance in total fibrinogen.

Total fibrinogen and IL-6 concentrations were generally higher than those reported in Europeans [13, 59]. IL-6 correlated positively with fibrinogen concentration, thereby accelerating clot formation and increasing fibre diameter, consequently contributing to clot pathology associated with CVD [68]. In the present study, higher IL-6 was associated with a faster CLT, independent of fibrinogen concentration and clot structure. This is probably the result of increases in other biomarkers influencing CLT, such as plasminogen leading to faster clot lysis. Previous reports have positively associated IL-6 with increased plasminogen transcription [69, 70].

Fibrinogen γ’ was also positively associated with IL-6, although the non-significant association of IL-6 with the % γ’ fibrinogen revealed the observation to be largely a reflection of the relationship with total fibrinogen. This finding suggests that in this study population, characterised by chronic low-grade inflammation, the influence of IL-6 on total and γ’ fibrinogen is probably due to up-regulation of the entire fibrinogen gene cluster, while alterations in the FGG alternative splicing mechanism may be more relevant in pronounced inflammatory conditions such as have been reported for CVD [26, 27].

IL-6-induced fibrinogen gene expression can be altered by polymorphic variation [16, 17]. Therefore, it is important to take genotype-IL-6 interactions into consideration when determining the influence of genetic variance on the fibrinogen phenotype, particularly in
populations with prevalent inflammatory conditions. Nine SNPs interacted significantly with IL-6 in predicting fibrinogen concentrations. These interactions furthermore led to altered fibrin clot properties, in particular increased fibre diameter, which has been associated with CVD [71]. The number of significant genotype-IL-6 interactions was reduced when individuals with high IL-6 concentrations (>100 pg/mL) were removed from analyses, suggesting that these interactions are physiologically more relevant in the presence of high IL-6 concentrations. The interaction with FGG-rs1049636 remained, however, indicating its potential relevance to fibrinogen regulation during chronic low-grade inflammatory conditions.

In addition, these interactions had an additive effect that could only be detected because of the lack of LD in the fibrinogen gene cluster in this African study population. The additive effect reached a threshold when more than three risk alleles occurred together. This observation should be investigated in a larger study population to allow analysis of the number of risk alleles, rather than just the presence thereof, thereby exploring dominant/co-dominant/recessive effects. Nonetheless, the observed additive effects reflect true physiological mechanisms, as the particular combinations are harboured concurrently in each individual. This data are in agreement with the findings of Ken-Dror et al. [63] who suggested that the fibrinogen phenotype is not regulated by one functional SNP only, but by a combination of minor alleles spanning the whole cluster. The lack of a causal contribution of fibrinogen to CVD as suggested by Mendelian randomisation studies may, therefore, need to be revisited, because these studies investigated single variants only and focussed on fibrinogen concentration alone while omitting qualitative effects such as altered clot properties [72]. Although our sample size was relatively small for genetic association studies, it did allow for the detection of significant results. Future studies should include a large study population from genetically diverse ethnic groups for replication and validation of current results.

In this Tswana population, of the 14 SNPs investigated, FGB-rs7439150, –1420G/A and –148C/T had the most pronounced effects on the fibrinogen phenotype in terms of both concentration and altered clot properties, with several of these associations altered by IL-6 concentrations. Two of these SNPs, FGB-rs7439150 and –1420G/A were also the top SNPs associated with fibrinogen concentrations in the largest fibrinogen GWAS to date ($\beta = 0.031, p = 9.5E^{-181}$; $\beta = 0.031, p = 6.9E^{-180}$, respectively) [61]. FGB –148C/T was not investigated most likely due to complete linkage with FGB –1420G/A in this European study population [61].

FGB-rs7439150 is located in a regulatory feature (ENSR00000175110) spanning 623 base pairs (4:154560017–154560640), and is predicted to have methylation potential, so could influence transcription [73]. In addition, possible functionality of –148C/T has been described by Verschuur et al. [17] owing to its ability to alter fibrinogen’s response to IL-6 by interfering with the hepatocyte nuclear factor 3 and CCAAT enhancer binding protein binding sites. Although the association of the FGB promoter –1420G/A polymorphism with fibrinogen concentration has been reported previously [9, 54, 61], it is not associated with any known regulatory features [73] and further investigation into its functionality is required. In this study population, these three SNPs were also in high, albeit not complete LD ($D’ > 0.92, r^2 > 0.82$), therefore, the possibility of lack of independence cannot be excluded. They did, however, have unrelated independent genotype-phenotype associations; FGB-rs7439150 with fibrinogen $\gamma$, FGB –1420G/A with total fibrinogen, and FGB –148C/T with maximum absorbance, respectively. Distinct differences in their IL-6-interactive associations (Tables 3 and 4) were also observed, strengthening their hypothesised independent functional contribution to the fibrinogen-related phenotypes. The possibility that other SNPs in strong LD with those we genotyped, were responsible for the associations observed, can of course not be ruled out. In addition there are many more (novel and known) SNPs in the fibrinogen gene cluster, which were not investigated, that may also influence the fibrinogen phenotype [9, 61, 67].
Conclusion

The genotypic exploration of the fibrinogen phenotype in the Tswana population showed the unique genetic composition of black South Africans even though this study did not attempt to characterise all the variation in the fibrinogen genes of this African population. It focused instead on SNPs previously indicated in the literature to have functional effects which could, to date, not be confirmed as a result of their high LD. It is clear that data obtained from European research cannot be extrapolated directly to Africans and that the risk profile in terms of the relative contribution of genetics and environmental factors might differ. Our results demonstrate that fibrinogen SNPs can modulate the influence of IL-6 on fibrinogen concentration individually and in an additive manner, with the presence of more minor alleles across these genes leading to greater increases in IL-6-induced fibrinogen expression in an apparently healthy African study population, albeit one with apparent chronic, low-grade inflammation. Therefore, in future, when investigating the effect of fibrinogen genetics on fibrinogen concentrations and CVD outcome, the possible interactions with modulating factors and the fact that SNP effects seem to be additive should be taken into account.

Supporting information

S1 Table. Primer and synthetic control sequences used for KASP analyses.
(DOCX)

S2 Table. Basic descriptive characteristics of the study population as published by Kotzé et al., (2015).
(DOCX)

S3 Table. Associations of individual SNPs with fibrinogen γ', % γ' and total fibrinogen as published by Kotzé et al., (2015).
(DOCX)

S4 Table. Associations of individual SNPs with clot-related phenotypes as published by Kotzé et al., (2015).
(DOCX)

Acknowledgments

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Author Contributions

Conceptualization: Fiona R. Green, Marlien Pieters.

Formal analysis: H. Toinét Cronjé.

Supervision: Marlien Pieters.

Writing – original draft: H. Toine Cronje, Marlien Pieters.

Writing – review & editing: H. Toine Cronje, Cornelie Nienaber-Rousseau, Lizelle Zandberg, Zelda de Lange, Fiona R. Green, Marlien Pieters.

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