A Genome-Wide Association Study Identifies UGT1A1 as a Regulator of Serum Cell-Free DNA in Young Adults: The Cardiovascular Risk in Young Finns Study

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Abstract

Introduction: Circulating cell-free DNA (cf-DNA) is a useful indicator of cell death, and it can also be used to predict outcomes in various clinical disorders. Several innate immune mechanisms are known to be involved in eliminating DNA and chromatin-related material as part of the inhibition of potentially harmful autoimmune responses. However, the exact molecular mechanism underlying the clearance of circulating cf-DNA is currently unclear.

Methods: To examine the mechanisms controlling serum levels of cf-DNA, we carried out a genome-wide association analysis (GWA) in a cohort of young adults (aged 24–39 years; n = 1841; 1018 women and 823 men) participating in the Cardiovascular Risk in Young Finns Study. Genotyping was performed with a custom-built Illumina Human 670 k BeadChip. The Quant-iT™ high sensitivity DNA assay was used to measure cf-DNA directly from serum.

Results: The results revealed that 110 single nucleotide polymorphisms (SNPs) were associated with serum cf-DNA with genome-wide significance (p<5×10^-8). All of these significant SNPs were localised to chromosome 2q37, near the UDP-glucuronosyltransferase 1 (UGT1) family locus, and the most significant SNPs localised within the UGT1 polypeptide A1 (UGT1A1) gene region.

Conclusion: The UGT1A1 enzyme catalyses the detoxification of several drugs and the turnover of many xenobiotic and endogenous compounds by glucuronidating its substrates. These data indicate that UGT1A1-associated processes are also involved in the regulation of serum cf-DNA concentrations.

Introduction

Circulating cell-free DNA (cf-DNA) was first described in 1940s [1], and it has frequently been used as a convenient indicator of cell death and tissue damage in various acute and chronic disorders [2–4]. Detectable amounts of cf-DNA are present in healthy individuals, yet significantly increased plasma/serum levels of cf-DNA have been reported in patients with acute cardiovascular disease, sepsis, cancer and pre-eclampsia [2–4]. Circulating nucleosomal DNA has also been proposed as a major autoantigen in systemic lupus erythematosus [5] and as a potential player in lupus nephritis [6]. Additionally, genomic DNA fragments released during sterile thyroid injury may also trigger autoreactivity-related thyroid dysfunction [7]. Other potentially detrimental functions implicated for cf-DNA include the horizontal transfer of oncogenic sequences (i.e. genometastasis) [8] and the creation of a proinflammatory milieu in the circulation [9]. Endogenous or microbial DNA are also known to act as alerting signals to the host immune system, and various soluble, membrane-bound and intracellular receptors that can recognise DNA have been
identified. These receptors include pattern recognition molecules such as collectins, ficolins, pentraxins, soluble CD14 and C1q [10]; Toll-like receptors (TLRs) [11]; the high-mobility group box such as collectins, ficolins, pentraxins, soluble CD14 and C1q [10]; and extrachromosomal histone H2B [7]. Upon sensing DNA, these receptors can mount an immuno-inflammatory response to eliminate the circulating DNA and prevent anti-DNA autoimmunity.

Despite its proven utility in diagnosis and in determining various clinical conditions, many aspects of its origin, metabolism and non-pathological fluctuations of cf-DNA remain unresolved. According to the current understanding, the majority of cf-DNA is derived from apoptotic or necrotic cells; however, active secretion by leukocytes may also contribute to the pool of circulating DNA [4]. Serum cf-DNA is hypothesised to consist of both free DNA and particle-bound forms [4], yet the relationships between cf-DNA composition, originating cell types and the given clinical conditions are unclear.

Studies in mice have demonstrated that injected DNA, including single-stranded DNA (ssDNA), double-stranded DNA (dsDNA), nucleosomal core particles and oligonucleotides, is cleared from the system very rapidly, in less than 40 minutes, and that the major organ responsible for the solubilisation and generation of DNA degradation products is the liver [14–16]. Nevertheless, the clearance kinetics of these distinct DNA species are somewhat different, and the induction of the acute phase response or co-injection of the acute phase reactants serum amyloid P (SAP) and C-reactive protein (CRP) causes a delay in chromatin clearance, concomitant with an increased localisation of cf-DNA to the liver [17]. However, it has also been demonstrated that macrophages play an essential role in processing the cf-DNA released from dead or dying cells [18,19]. Data regarding the corresponding cf-DNA dynamics in humans are very scarce, but it has been reported that foetal cf-DNA has a mean half-life of 16.3 minutes in the maternal plasma and that plasma nucleosomes have only a partial role in the removal of foetal DNA [20].

To identify the genetic determinants of serum cf-DNA levels and thus elucidate the mechanisms involved in cf-DNA turnover, we performed a genome-wide association analysis (GWA) in a cohort of young adults participating in the Cardiovascular Risk of Young Finns study.

Results

The characteristics of the study population are presented in Table 1. In addition to age and insulin concentration, all of the variables tested differed significantly between women and men (Table 1). In the GWA, a total of 110 single nucleotide polymorphisms (SNPs) were associated with serum cf-DNA levels with a genome wide significance level of p < 5×10^{-8}. All 110 SNPs localised to the *UGT1* polygene cluster of the *UGT1A1* (UGT1) family locus on chromosome 2q37, and the most significant SNPs resided in the *UGT1A1* polygene 1 (UGT1A1) gene region (Figure 1 and Figure 2).

In the *UGT1A1* gene family region, we observed a major haploblock that could be further divided into nine sub-blocks (using the four gamete rule in HaploView). The four top SNPs on *UGT1A1* belong to the sub-block no. 8, which spans approximately 11 kb, and these SNPs appeared to be in virtually perfect linkage (r^2 = 0.9991 − 1.0). None of the significant SNPs in the sub-block no. 8 resided in exonic regions. However, among the significant SNPs outside this haploblock, three non-synonymous SNPs (rs6759892, rs2070959, rs1105879, p = 8.2×10^{-10} − 3.4×10^{-14}) were identified in the coding region of *UGT1A6* (Figure 2). In addition, three significant SNPs (rs17864701, rs6663967, rs6741669, p = 1.2×10^{-21} − 1.1×10^{-17}) were detected in the

### Table 1. Characteristics of the study population.

| Variable                  | Women (n = 1018) |       | Men (n = 823) |       | p for difference |
|---------------------------|------------------|-------|--------------|-------|-----------------|
| Age (years)               | 31.7             | 5.0   | 31.7         | 5.0   | N.S.            |
| BMI (kg/m^2)              | 24.4             | 4.5   | 25.7         | 4.2   | <0.001          |
| Waist circumference (cm)  | 79.2             | 11.2  | 90.0         | 11.0  | <0.001          |
| Systolic blood pressure (mmHg) | 113               | 12    | 121          | 13    | <0.001          |
| Diastolic blood pressure (mmHg) | 69              | 10    | 73           | 11    | <0.001          |
| Total cholesterol (mmol/L) | 5.02             | 0.87  | 5.20         | 1.00  | <0.001          |
| HDL-cholesterol (mmol/L)  | 1.39             | 0.29  | 1.16         | 0.28  | <0.001          |
| LDL-cholesterol (mmol/L)  | 3.11             | 0.76  | 3.40         | 0.89  | <0.001          |
| Triglycerides (mmol/L)*   | 1.0              | 0.8–1.3 | 1.2        | 0.9–1.8 | <0.001        |
| Insulin (mU/L)*           | 6                | 5–9   | 6            | 4–9   | N.S.            |
| Glucose (mmol/L)*         | 4.9              | 4.6–5.1 | 5.1        | 4.9–5.4 | <0.001        |
| Homocysteine (µmol/L)*    | 8.5              | 7.2–10.1 | 10.1       | 8.7–11.7 | <0.001       |
| cf-DNA (µg/ml)            | 0.755            | 0.135 | 0.862        | 0.111 | <0.001          |
| CRP (mg/L)*               | 0.84             | 0.35–2.30 | 0.59       | 0.30–1.42 | <0.001     |
| Alcohol (drinks per week)*| 2                | 0–5   | 6            | 1–13  | <0.001          |
| Smoking daily (% of total)† | 19.3            |       | 29.4         |       | <0.001          |
| Use of COCs (% of total)  | 24.9             |       |              |       |                 |

*Median values and interquartile range (IQR), Mann-Whitney test for between-group differences.
†Chi-squared test for differences between groups.

Abbreviations: BMI, body mass index; cf-DNA, cell-free DNA; COCs, combined oral contraceptives; CRP, C-reactive protein; N.S., not significant.

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promoter region of DNAJB3 (Figure 2). The top SNPs in each of the nine sub-blocks are presented in Table 2.

When the selected tag SNPs from each sub-block were analysed using the stepwise AIC model selection algorithm, only the tag SNP rs4148324 which identifies the sub-haploblock no. 8 containing the four top SNPs, remained in the model. This SNP was found to explain 5.3% of the total variation in serum cf-DNA levels. In addition, conditioning the analysis on the top SNP rs4148324 (i.e. additional adjustment of the model with the top SNP) did not reveal any other SNPs associated with cf-DNA levels at a genome-wide significance level (all p-values > 0.04).

An interaction network for UGT1A1 constructed with IPA (Ingenuity Pathway Analysis) revealed that several molecules in the UGT1A1 interaction network are connected to DNA metabolism, quantity, fragmentation and synthesis of DNA (Figure 3). In addition, molecular connections to HMGB signalling and TLR signalling were observed for UGT1A1 in the IPA Canonical Pathway analysis (Figure 3).

Discussion

The circulating cf-DNA concentration has been shown to increase during pathological processes, reflecting the rate of cellular death and tissue damage. However, the regulation and metabolism of cf-DNA have received much less attention, and data regarding the turnover of cf-DNA are scarce. In this study, we demonstrate that a novel candidate gene, UGT1A1, may contribute to the regulation of serum cf-DNA in young adults. The UGTs comprise a family of enzymes that primarily transfer glucuronic acid to hydroxyl, carboxylic acid and amine group-bearing compounds to facilitate the catabolism and clearance of these compounds [21]. Because a wide variety of compounds contain these groups, glucuronidation provides a common mechanism for the elimination of several endobiotics, xenobiotics and drugs. The UGT1A1 locus encodes several isoforms of the enzyme, each with a tissue-specific expression pattern [21]. Although the liver is the major producer of UGT1A1, the expression of this isoform has also been detected in the bile duct, the stomach, the small intestine, the kidneys, the bladder, the uterus and the colon [21,22]. Interestingly, the expression of UGT1A1 has recently been demonstrated in human peripheral blood mononuclear cells [23].

Due to the strong linkage disequilibrium between the SNPs in the UGT1 region (Figure 2) we cannot completely rule out the involvement of the other significant SNPs, such as those in the coding region of UGT1A6 or in the promoter of DNAJB3, in the regulation of cf-DNA levels. Nevertheless, the main signal can be confined to the sub-block of the four top UGT1A1 SNPs because the significance of the p-values of the other SNPs decreases along with the increase in distance from this sub-block (Table 2 and Figure 2).

One of the best characterised functions of UGT1A1 is the glucuronidation of bilirubin [21], and numerous recent GWA studies have identified UGT1A1 alleles as the major regulators of serum bilirubin levels [24]. The four most significant SNPs associated with serum cf-DNA levels in this study (rs4148324, rs6742078, rs4148325 and rs887829) have also been identified as regulators of serum bilirubin concentrations [24]. The effect of these SNPs on bilirubin levels is, however, reported to be due to a functional TA-repeat polymorphism in the UGT1A1 promoter TATA-box area; recent conditional linkage scans of this TA-repeat association demonstrated that it accounts for the linkage peaks observed for the other associated loci in the gene region [25,26]. Given that the ancestral alleles of the rs4148324, rs6742078, rs4148325 and rs887829 SNPs are in strong linkage with the wild-type form of the TA-repeat polymorphism (TA6, rs8175347), and because all of these variants belong to the same haploblock in our population, we assume that the same regulation pattern holds true for both serum cf-DNA and bilirubin levels. The TA6 form of the repeat polymorphism is associated with lower bilirubin levels due to higher promoter activity (via optimal transcription factor IID binding), resulting in higher UGT1A1 enzyme levels and enhanced bilirubin glucuronidation and
Because we observed lower cf-DNA levels in subjects bearing the wild type alleles ($\beta = -0.3358$) of the four top SNPs in comparison with the minor allele carriers, we suggest that higher UGT1A1 production, or some cellular process associated with UGT1A1 induction, contributes to more effective turnover of serum cf-DNA.

Animal studies have demonstrated that the liver is the major site of cf-DNA clearance, although some portion of the circulating DNA also localises to other organs, such as the kidneys and spleen [14,16,17]. However, in humans, foetal cf-DNA has been shown to be cleared rapidly from the maternal circulation, and plasma nucleases have only a partial effect on cf-DNA disposal [20]. Moreover, relatively large DNA fragments (>150 base pairs) originating from non-renal or urinary tract tissues have been detected in the urine [28], indicating that the complete digestion of cf-DNA by nucleases is not required for its excretion. Beyond these observations, the metabolism and fates of different cf-DNA forms are unknown. Although our data does not provide a mechanistic basis of UGT1A1-mediated cf-DNA clearance, we propose that the metabolism of cf-DNA may be facilitated by UGT1A1, either directly by glucuronidation or indirectly via the interactions between UGT1A1 and the molecules related to DNA sensing and processing (Figure 3). It is also possible that a portion of the cf-DNA is endocytosed and/or co-processed with the traditional UGT1A1 substrates, such as the bilirubin-albumin complex [29].

Nevertheless, physiological interactions between UGT1A1 and nucleotides have been demonstrated [30] and the glucuronidation of a 2',3'-dideoxynucleoside compound, the 3'-Azido-3'-deox-
ythymidine (AZT), has been demonstrated, although this reaction is performed by the UGT2B7 enzyme [31]. However, at the present time, we can only speculate whether these observations are related to the glucuronidation of endogenous cf-DNA during the course of its cellular internalisation or (apoptotic) release. Neither can we exclude the possibility that the actual UGT1A1 substrate might be some lipophilic or proteinaceous structure associated with cf-DNA. Furthermore, the potential involvement of extrahe-

Table 2. The top SNPs in each of the 9 sub-blocks in the UGT1A gene region (see Figure 2).

| SNP   | Sub-block | Locus (bp) | Ancestral allele | Minor allele | MAF (b) | MAF (S.E.) | p* | Imputed | Location |
|-------|-----------|------------|------------------|--------------|---------|------------|----|---------|----------|
| rs17864683 | 1          | 234243948   | A                | C            | 0.3221  | 0.3246 (0.0352) | 2.06E-14 | yes      | intron, UGT1A10 |
| rs10168416  | 2          | 234261826   | G                | C            | 0.4230  | 0.3236 (0.0325) | 2.98E-15 | no       | intron, UGT1A8   |
| rs1105879   | 3          | 234266941   | C                | A            | 0.4603  | 0.3127 (0.0324) | 8.20E-18 | no       | missense, UGT1A6 |
| rs17863787  | 4          | 234275833   | G                | T            | 0.3592  | 0.3092 (0.0338) | 1.40E-19 | no       | intron, UGT1A6   |
| rs6744284   | 5          | 234290036   | T                | C            | 0.3710  | 0.2618 (0.0335) | 3.04E-20 | no       | intron, UGT1A6   |
| rs672076    | 6          | 234312056   | A                | G            | 0.3705  | 0.2824 (0.0334) | 1.25E-21 | yes      | intron, UGT1A6   |
| rs17864701  | 7          | 234317456   | T                | C            | 0.3722  | 0.2591 (0.0335) | 1.19E-21 | yes      | intron, UGT1A6   |
| rs887829    | 8          | 234333309   | A                | G            | 0.3983  | 0.3358 (0.0331) | 1.59E-23 | no       | intron, UGT1A1   |
| rs6742078   | 9          | 234337378   | T                | G            | 0.3983  | 0.3358 (0.0331) | 1.59E-23 | no       | intron, UGT1A1   |
| rs4148324   | 10         | 234337461   | G                | T            | 0.3983  | 0.3358 (0.0331) | 1.59E-23 | no       | intron, UGT1A1   |
| rs4148325   | 11         | 234338048   | T                | C            | 0.3983  | 0.3358 (0.0331) | 1.59E-23 | no       | intron, UGT1A1   |
| rs11690786  | 12         | 234357356   | T                | C            | 0.3839  | 0.2711 (0.0337) | 9.67E-17 | yes      | intron, HEATR7B1 |

The four SNPs that are associated most significantly with serum cf-DNA levels are in sub-block 8.

*For each SNP, the ancestral allele was modelled, and the β coefficient represents the change in cf-DNA level (μg/ml) with one additional copy of the ancestral allele.

*Adjusted for age, sex, BMI and genetic East-West stratification in the Finnish population.

Abbreviations: SNP, single nucleotide polymorphism; Chr, chromosome; bp, base pairs; MAF, minor allele frequency; S.E., standard error.

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Figure 3. The UGT1A1 interaction network generated using the Ingenuity Pathway Analysis. The molecular relationships connecting UGT1A1 to DNA metabolism synthesis, fragmentation and quantity of DNA are indicated with Fx symbols, and the associated Canonical Pathways are indicated with CP symbols. Abbreviations: AHR, aryl hydrocarbon receptor; HMGB1, high-mobility group box protein 1; IL1B, interleukin-1 beta; MYC, v-myc myelocytomatosis viral oncogene homolog; NR113, nuclear receptor subfamily 1, group I, member 3; PPARC1A, peroxisome proliferator – activated receptor gamma coactivator 1 alpha; RELA, v-rel reticuloendotheliosis viral oncogene homolog A (avian); TLR4, Toll-like receptor 4.

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patic UGT1A1-expression sites, such as leukocytes [23] or intestinal cells [32], in cf-DNA turnover cannot be addressed by our data.

Our statistical model for genetic variation in the UGTA1 explains only 5.3 % of the total variation in cf-DNA levels, although the association is highly statistically significant (p = 1.6 x 10^-55). The somewhat modest coefficient of determination can be explained by the complex nature of the serum cf-DNA; different forms of cf-DNA might be assigned to different clearance pathways with dissimilar kinetics and saturability. It is also known that the serum contains higher cf-DNA levels than the plasma, allegedly due to the release of genomic DNA from leukocytes during blood clotting [33] Recently, however, this view has been challenged by the observation that leukocyte rupture is not the major factor causing increased serum cf-DNA levels [34]. Nevertheless, the strong association suggests a significant biological role for UGT1A1 in regulating cf-DNA levels, especially because no other polymorphisms outside the reported UGT1A1 loci emerged with genome-wide significance when the analysis was adjusted using the top SNP (rs4148324).

In conclusion, the results of this GWA study demonstrate that UGT1A1 polymorphisms are associated with serum cf-DNA levels in young Finns. We propose that UGT1A1-associated processes are, either directly or indirectly, involved in the regulation of cf-DNA concentration. These results, however, are limited to one study cohort which is a major limitation of our study. Therefore the results should be replicated in an independent study cohort, and further investigation to establish the exact role of UGT1A1 in cf-DNA turnover is warranted.

**Materials and Methods**

**Study population**

The study population consisted of 1841 participants in the Cardiovascular Risk in Young Finns study (aged 24–39 years; 1018 women and 823 men). The study was approved by the local ethics committees (the University Hospitals of Helsinki, Turku, Tampere, Kuopio and Oulu) and was conducted following the guidelines of the Declaration of Helsinki. All participants gave their written informed consent. For a detailed cohort description and the assessment of the variables listed in Table 1, see [35] and the references therein. The data in this study were from individuals with successful cf-DNA measurements and genotyping and who participated in the follow-up study in 2001.

**Quantification of serum cf-DNA**

Total circulating cf-DNA was determined directly in previously thawed serum using the fluorescence-based Quant-iT™ high-sensitivity DNA assay kit and a Qubit® fluorometer (Invitrogen, Carlsbad, CA, USA). All samples were analysed in duplicate, and the mean of the two measurements was used as the final value. At the mean serum cf-DNA levels of 0.650 µg/ml, the assessed intra- and inter-day variation coefficients were 2.2% and 4.7%, respectively. At the mean serum cf-DNA level of 1.02 µg/ml, the intra- and inter-day variation coefficients were 3.0% and 5.6%, respectively.

**Genotyping**

Genotyping was performed at the Welcome Trust Sanger Institute using a custom made Illumina Human 670 k BeadChips. Genotypes were determined using the Illumina clustering algorithm [36]. Fifty-six samples failed the Sanger genotyping pipeline QC criteria (i.e., duplicated samples, heterozygosity, low call rate, or Sequenom fingerprint discrepancies). Three samples were removed due to a low genotyping call rate (<0.95) and 54 samples were excluded for possible relatedness (pi.hat>0.2). A total of 11,766 single SNPs were excluded due to deviation from Hardy-Weinberg equilibrium (HWE) test (p<1e-06). 7,746 SNPs failed the missingness test (call rate<0.95) and 34,596 SNPs failed the frequency test (MAF<0.01). After quality control, 546,677 genotyped SNPs remained available for further analysis. Genotype imputation was performed using MACH 1.0 [37,38] and HapMap II CEU (release 22) samples as reference. After imputation, 2,543,887 imputed SNPs were available. SNPs with squared correlations ≥0.3 between imputed and true genotypes were considered well imputed.

**Statistical methods**

The comparison of the basic study population variables presented in Table 1 was conducted with the Student’s t-test, Mann-Whitney’s test or chi-squared test. Prior to the GWA, all continuous variables were subjected to inverse normal transformation to minimise the incidence of type I errors and to reduce the impact of outliers [39]. A stepwise backward model (Akaike information criterion, AIC) was used to determine which covariates explained the most of the variation in the cf-DNA levels. In addition to the variables listed in Table 1, genetic main identity-by-descent (IBD) clustering components and interactions between age and gender as well as smoking status and the use of alcohol were included in the model. The variables that remained in the model and were thus used as the adjustment covariates in GWA were gender, age, systolic blood pressure, fasting glucose, triglycerides, CRP, homocysteine, daily smoking, use of alcohol, use of combined oral contraceptives, main IBD components and the gender*age and daily smoking*use of alcohol interaction terms. These factors were observed to account for 30.5% of the variation in cf-DNA levels. To reveal other SNPs potentially associated with serum cf-DNA, the analysis was additionally adjusted with the top SNP, rs4148324. Standardised residuals were extracted from the model, and association analysis was performed using linear regression with an assumption of an additive genetic effect. PLINK [40] was used for the true genotyped SNPs, and ProbaBEL [41] was used for imputed genotype dosages. If the same SNP was available in both genotyped and imputed form, the genotyped form was displayed and included in the results. A commonly accepted threshold for genome-wide statistical significance level (p<5 x 10^-8) was used to identify significant SNPs. Manhattan and Q-Q plots were drawn to confirm the validity of the analysis. The genomic inflation factor (lambda) was 0.9996475. The variance in serum cf-DNA level explained by each SNP (R^2) was calculated as follows: R^2 = 1/(1+[number of samples-2]*SE of SNP/beta estimate of SNP^2]). The individual genotype data for genome-wide significant SNPs in the same region were extracted from both the genotyped and imputed data; then the allele dosages of the imputed SNPs were rounded to best guess genotypes and transferred to Haploview [42] for haploblock analysis and tag SNP identification. Haploblocks were defined using the four gamete rule in Haploview, and the SNP with the lowest p-value in each haploblock was selected as the tag SNP. All tag SNPs were analysed with stepwise backward AIC to identify the potential independent signals in different haploblocks. Unless otherwise noted, the statistical analyses were performed the appropriate packages in R (MASS for stepwise AIC).

To identify the potential interactions via which UGT1A1 could be connected to cf-DNA regulation, a graph of the UGT1A1 molecular network and associated functions and Canonical Pathways was generated using IPA (Ingenuity® Systems, www.
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Author Contributions

Performed the experiments: JJ LPL. Analyzed the data: JJ LPL. JK. Contributed reagents/materials/analysis tools: TL MH JV OTR MK NHK. Wrote the paper: JJ LPL.

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