Overlapping genetic susceptibility variants between three autoimmune disorders: rheumatoid arthritis, type 1 diabetes and coeliac disease

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Abstract

Introduction: Genome wide association studies, replicated by numerous well powered validation studies, have revealed a large number of loci likely to play a role in susceptibility to many multifactorial diseases. It is now well established that some of these loci are shared between diseases with similar aetiology. For example, a number of autoimmune diseases have been associated with variants in the PTPN22, TNFAIP3 and CTLA4 genes. Here we have attempted to define overlapping genetic variants between rheumatoid arthritis (RA), type 1 diabetes (T1D) and coeliac disease (CeD).

Methods: We selected eight SNPs previously identified as being associated with CeD and six T1D-associated SNPs for validation in a sample of 3,962 RA patients and 3,531 controls. Genotyping was performed using the Sequenom MassArray platform and comparison of genotype and allele frequencies between cases and controls was undertaken. A trend test P-value < 0.004 was regarded as significant.

Results: We found statistically significant evidence for association of the TAGAP locus with RA (P = 5.0 × 10^-4). A marker at one other locus, C1QTNF6, previously associated with T1D, showed nominal association with RA in the current study but did not remain statistically significant at the corrected threshold.

Conclusions: In exploring the overlap between T1D, CeD and RA, there is strong evidence that variation within the TAGAP gene is associated with all three autoimmune diseases. Interestingly a number of loci appear to be specific to one of the three diseases currently studied suggesting that they may play a role in determining the particular autoimmune phenotype at presentation.

Introduction

Rheumatoid arthritis (RA) is an auto-immune, inflammatory joint condition, affecting 0.8% of the UK population [1]. It has long been established that the major genetic predisposition to this disease is contributed by variants of the class II HLA gene, HLA DRB1. Both type 1 diabetes (T1D) and coeliac disease (CeD) are also auto-immune, inflammatory diseases for which the major genetic contribution arises from the major histocompatibility complex and which are also characterised by autoantibody formation [2].

Recent genome wide association and subsequent replication studies in these three diseases have revealed a large number of well validated, non-HLA genetic risk loci providing an opportunity to explore the possibility of overlapping susceptibility between them [3]. Testing confirmed T1D susceptibility loci in CeD samples has already resulted in the identification of novel CeD loci [2].
A previous study exploring the overlap of CeD and RA susceptibility loci identified association to six regions [4]. We have recently reported our findings of the testing of putative T1D susceptibility loci in RA, which identified AFF3 as a novel RA susceptibility locus [5]. However, exploring the overlap between the three diseases in more detail may provide a further opportunity to elucidate the genetic similarities, and differences, between these common autoimmune diseases.

We, therefore, tested 14 validated susceptibility loci from CeD and T1D in a large, well powered cohort of British Caucasian patients with RA and independent controls to explore the overlap between loci identified in all three conditions.

Materials and methods

Samples

Clinical characteristics of the RA case-control cohort tested have been described previously [6]. RA cases satisfied American College of Rheumatology (ACR) classification criteria for RA modified for genetic studies, all provided informed consent and were collected with ethical committee approval (North-West Multi-Centre Research Ethics Committee (MREC 99/8/84) and the University of Manchester Committee on the Ethics of Research on Human Beings) [7,8]. Their clinical characteristics are shown in Table S1 in Additional file 1.

SNP selection

SNP markers with evidence for association with CeD susceptibility ($P < 1 \times 10^{-5}$) in reference [2] were selected for genotyping in a cohort of RA and control samples. Where more than one SNP was associated at a particular locus with CeD, the most associated variant was selected for genotyping in the RA case-control cohort.

SNP markers with evidence for association in T1D in the same reference, but not genotyped as part of our previous work, were also selected for genotyping in the current cohort of RA cases and controls.

Genotyping

SNP markers were genotyped using the iPlex chemistry on the Sequenom platform according to the manufacturer’s instructions (Sequenom, San Diego, California, USA) [9]. Duplicate samples and negative controls were included on the plates to ensure genotyping accuracy. Additional quality control measures were introduced such that only SNP markers and samples exceeding a 90% genotyping success threshold were included in any subsequent analysis.

Analysis

The power to detect association in the current cohort assuming the same odds ratio (OR) as was reported in the CeD disease study was calculated using Quanto [10]. Departure from Hardy-Weinberg equilibrium was tested for all SNP markers. Genotype frequencies were compared between RA cases and controls using the trend test implemented in STATA (StataCorp LP, Texas, USA). Conditional logistic regression was used to assess whether effects were independent where two SNPs had been tested at the same locus.

Results

SNP selection

Eight CeD SNPs were selected for genotyping as they had not previously been examined for association with RA (Table 1). We have previously reported our results for most but not all of the T1D loci tested in a recent study exploring the genetic overlap between CeD and T1D [5]. For completeness, six T1D associated SNPs not previously investigated for association with RA in the current data set were, therefore, also tested (Table 2).

Genotyping

Genotyping was undertaken in an available validation cohort of 3,962 RA cases and 3,531 controls from the UK. A $P$-value of 0.004 was selected as the nominal significance threshold. The study had between 67% and 99% power to detect association with the CeD loci in this sample at the $P < 0.004$ threshold assuming the same allele frequencies and effect sizes as reported with CeD (Table S2 in Additional file 1). Duplicate samples showed >99% concordance. All samples and SNPs met the quality control threshold of >90% genotyping success. All SNP genotype frequencies conformed to Hardy-Weinberg expectations in the controls.

One marker with previous evidence for association with CeD was also associated with RA in the current cohort: rs182429 mapping to the TAGAP gene OR 0.88, 95% confidence intervals (CI) 0.82 to 0.95; $P$-trend 0.0005) (Table 1). The TAGAP locus has previously been identified as an RA susceptibility locus following meta-analysis of genome-wide association studies in RA followed by validation in independent samples, which included samples from the current study [11]. However, a different SNP marker (rs394581) was tested in that study. The two SNPs were in moderate linkage disequilibrium (LD) ($r^2 = 0.32; D' = 0.73$) in the control samples tested here but conditional logistic regression analysis suggested that the association arises primarily from the CeD SNP, rs182429 ($P$-value for rs394581 after conditioning on rs182429 = 0.90).

Of the T1D SNPs tested, only rs229541, mapping to CIQTNF6, showed nominal evidence for association but this did not retain statistical significance at the threshold selected (OR 1.08, 95% CI 1.0 to 1.15, $P = 0.04$) (Table 2).
| SNP      | CHR | position | Gene   | MAF cases | MAF controls | P_HWE | case_11    | case_12    | case_22    | control_11 | control_12 | control_22 | P_trend | Allelic OR (95% CI) |
|----------|-----|----------|--------|-----------|--------------|--------|------------|------------|------------|-------------|------------|-------------|----------|----------------------|
| rs2816316| 1   | 190803436|RGS1   | 0.17      | 0.18         | 0.31   | 119        | 1027       | 2501       | 99          | 835        | 1993        | 0.65     | 0.98 (0.89 to 1.07)  |
| rs917997 | 2   | 102437000|IL18RAP | 0.23      | 0.22         | 0.87   | 208        | 1,285      | 2,154      | 147         | 1,013      | 1,771       | 0.17     | 1.06 (0.98 to 1.15)  |
| rs6441961| 3   | 46327388 |CCR3    | 0.31      | 0.31         | 0.76   | 350        | 1,582      | 1,720      | 280         | 1,239      | 1,409       | 0.52     | 1.03 (0.95 to 1.1)   |
| rs333    | 3   | 46389951 |CCR5    | 0.11      | 0.11         | 0.38   | 23         | 689        | 2,802      | 35          | 642        | 2,444       | 0.08     | 0.91 (0.81 to 1.01)  |
| rs17810546| 3   | 161147744|IL12A   | 0.12      | 0.12         | 0.66   | 52         | 736        | 2,862      | 45          | 616        | 2,264       | 0.32     | 0.95 (0.85 to 1.05)  |
| rs1464510| 3   | 189595248|LPP     | 0.45      | 0.44         | 0.72   | 697        | 1,730      | 1,077      | 617         | 1,524      | 966         | 0.82     | 1.01 (0.94 to 1.08)  |
| rs182429 | 6   | 159389562|TAGAP   | 0.42      | 0.45         | 0.82   | 600        | 1,770      | 1,159      | 590         | 1,421      | 870         | 0.0005   | 0.88 (0.82 to 0.95)  |
| rs653178#| 12  | 110492139|SH2B3   | 0.49      | 0.51         | 0.68   | 914        | 1,744      | 985        | 771         | 1,449      | 703         | 0.02     | 0.92 (0.86 to 0.98)  |

*rs = 1 with rs3184504; 11 = homozygote for minor allele, 12 = heterozygote, 22 = homozygote for major allele; CHR, chromosome; MAF, minor allele frequency; P_HWE, P-value for Hardy Weinberg equilibrium test.*

| SNP      | CHR | position | Gene   | MAF cases | MAF controls | P_HWE | case_11    | case_12    | case_22    | control_11 | control_12 | control_22 | P_trend | Allelic OR (95% CI) |
|----------|-----|----------|--------|-----------|--------------|--------|------------|------------|------------|-------------|------------|-------------|----------|----------------------|
| rs11755527| 6   | 91014952 | BACH2  | 0.46      | 0.46         | 0.91   | 729        | 1,790      | 990        | 669         | 1,551      | 891         | 0.86     | 0.99 (0.93 to 1.06)  |
| rs689    | 11  | 21388000 | INS    | 0.28      | 0.28         | 0.54   | 287        | 1,411      | 1,818      | 254         | 1,246      | 1,612       | 0.95     | 1.00 (0.93 to 1.08)  |
| rs12708716| 16  | 11087374 | CLEC16A| 0.36      | 0.36         | 0.63   | 507        | 1,791      | 1,643      | 453         | 1,596      | 1,455       | 0.88     | 1.00 (0.93 to 1.06)  |
| rs3825932| 15  | 77022501 | CTSH   | 0.32      | 0.32         | 0.77   | 368        | 1,506      | 1,619      | 323         | 1,366      | 1,407       | 0.62     | 0.98 (0.91 to 1.06)  |
| rs3780013| 21  | 42714397 | UBASH3A| 0.45      | 0.43         | 0.86   | 705        | 1,752      | 1,052      | 590         | 1,522      | 996         | 0.07     | 1.07 (1.00 to 1.14)  |
| rs229541 | 22  | 35921264 | C1QTNF6| 0.45      | 0.43         | 0.49   | 726        | 1,693      | 1,100      | 564         | 1,548      | 1,007       | 0.04     | 1.08 (1.00 to 1.15)  |

*11 = homozygote for minor allele, 12 = heterozygote, 22 = homozygote for major allele; CHR, chromosome; MAF, minor allele frequency; P_HWE, P-value for Hardy Weinberg equilibrium test.*
Interestingly, this gene lies close to the IL2RB gene, which has previously been associated with RA susceptibility [6]. The apparent lack of association detected with T1D susceptibility loci in our sample may result from limited power to detect the effect sizes reported with the larger T1D samples: for three of the six loci, there was <40% power to detect association at the $P < 0.004$ threshold (Additional file 1 Table S2). However, association with the INS, CLEC16A and CTSH loci can be excluded with more confidence.

The three autoimmune diseases examined in the current study are characterised by the presence of autoantibodies and, interestingly, the association of the TAGAP variant is stronger in the subgroup of RA patients with anti-cyclic citrullinated peptide (anti-CCP) antibodies ($P$-trend $= 0.0001$, allelic OR 0.83 95% CI 0.75 to 0.91), suggesting a possible role in control of B cell function.

**Discussion**

By investigating the genetic overlap of CeD genes with RA, we have identified the TAGAP locus as being associated with RA. Our findings confirm and refine the results of a previous study in RA that also showed association to the TAGAP region but with a different variant; our results show that the same SNP that is associated with CeD and T1D also shows stronger evidence for association with RA [11]. Data from the current study extend that of previous work identifying that CTLA-4, the IL2_21 region, 6q23 (TNFAIP3), SH2B3, PRKCQ, MMELI and now TAGAP are susceptibility loci that are common to the three autoimmune diseases examined in the current study: T1D, CeD and RA [2,5].

The current study is limited by the fact that, because the identification of novel susceptibility markers is progressing rapidly, a number of additional CeD and T1D susceptibility loci have been identified more recently, which have not been tested as yet for association with RA [12-14]. Hence, this is not a comprehensive examination of the overlap among all T1D, CeD and RA loci. However, of the loci tested so far, there appears to be greater overlap between T1D/CeD and T1D/RA than between CeD and RA. This may reflect the fact that larger sample sizes were used to investigate T1D (> 8,000 T1D cases vs >9,000 controls) than CeD (2,500 cases vs >9,000 controls), thereby reducing the possibility of false negative results in the T1D series. The lack of association of some of the CeD loci with RA could reflect lack of power, particularly if *winner’s curse* has resulted in an overestimation of the effect sizes originally reported in the CeD studies. However, effect sizes observed when data from four GWAS of CeD were combined by meta-analysis were similar to those reported in the study by Smyth et al., which was used to select the CeD variants for testing in the current study [2,13]. That suggests that *winner’s curse* and over-estimation of effects sizes in CeD cannot explain the lack of association observed with RA.

The TAGAP minor allele confers protection against RA, similar to previous reports of T1D but contrasting with CeD in which the minor allele is associated with risk. Our results confirm and extend the evidence that there are common autoimmune susceptibility genes but suggest that the overlap of RA is stronger with T1D than CeD.

Relatively little is known about the TAGAP gene, which encodes a protein (T-cell activation RhoGTPase activating protein) transiently expressed in activated T cells, suggesting that it may have a role in immune regulation [2].

The TAGAP gene has previously been associated with RA following meta-analysis of US and European RA cohorts, although a different variant mapping to the same locus was genotyped [11]. Nonetheless, the same UK samples tested as part of the current study were also included in the validation phase of the meta-analysis and so data for both variants was available. The results of conditional logistic regression analysis suggest that the primary effect is with the CeD variant tested in the current study and that the association reported previously with rs394581, may have arisen due to LD, at least in the UK samples.

It is interesting to note that the SNP tested at the CCR5 locus which is associated with both T1D and CeD, also showed a trend towards association with RA. A previous meta-analysis of studies in RA investigating this deletion variant within the gene is suggestive of association with RA [15]. A combined meta-analysis, including data from the previous meta-analysis, two subsequent reports [16] and the current data, reveals statistically significant evidence for association ($P = 1.4 \times 10^{-5}$) but because of heterogeneity between the studies, these results should be interpreted with caution (Figure 1).

Exploring the genetic overlap between related diseases may reveal key common pathways that could suggest therapeutic targets applicable to more than one disease. However, the susceptibility loci unique to a particular disease are also of interest; differences may reflect genuine specificity between the diseases and may influence what determines the particular autoimmune phenotype. Of the confirmed non-HLA RA susceptibility loci (reviewed in [17]), only the CD40 and TRAF1/C5 variants have not been reported to be associated with either CeD or T1D although CD40 is associated with autoimmune thyroiditis [18]. Even the TRAF1/C5 association is not unique to RA as there have been reports of association with juvenile idiopathic arthritis and systemic lupus erythematosus [19,20]. Despite the inherent bias arising from the fact that loci associated with one autoimmune disease are often subsequently targeted for investigation in other autoimmune diseases, the degree of genetic
overlap is remarkable. One possible explanation may be that specificity of the autoimmune phenotype is determined by the particular HLA associations of each disease that could either directly or indirectly influence the response to environmental agents. This hypothesis would require further investigation in well-powered cohorts with both genetic and reliable environmental data.

In summary, we report association of RA with the TAGAP gene, identified through targeting loci previously associated with CeD. The findings identify that the same SNP associated with T1D and CeD shows stronger association with RA than a previously reported variant and extend the evidence for overlap between autoimmunity genes.

Conclusions
The TAGAP gene, previously associated with both T1D and CeD, is also associated with RA susceptibility. This confirms the finding of a previous study showing association of the TAGAP locus with RA but suggests that the variant associated with autoimmune diseases is more strongly associated than that reported previously.

Additional material

Additional file 1: Overlapping genetic susceptibility variants between three autoimmune disorders - supplementary information. The file contains tables showing the clinical characteristics of the RA patient samples tested as well as the power of the study for each SNP tested.

Abbreviations
AFF3: AF4/FMR2 family member 3; C1QTNF6: C1q and tumour necrosis factor related protein 6; CCR5: cytokine-chemokine receptor 5; CD40: CD40 molecule; CED: coeliac disease; CHR: chromosome; CI: confidence interval; CLEC16A: C-type lectin domain family 16; CTLA4: cytotoxic T lymphocyte activated 4; CTSH: cathepsin H; HLA: human leucocyte antigen; HWE: Hardy Weinberg equilibrium; IL: interleukin; IL2RB: interleukin 2 receptor beta; INS: insulin; LD: linkage disequilibrium; MAF: minor allele frequency; OR: odds ratio; PTPN22: protein tyrosine phosphatase non-receptor 22; RA: rheumatoid arthritis; SNP: single nucleotide polymorphism; T1D: type 1 diabetes; TAGAP: T-cell activation RhoGTPase activating protein; TNFAIP3: tumour necrosis factor activation induced protein 3; TRAF1/C5: tumour necrosis receptor activation factor 5/Complement 5; WTCCC: Wellcome Trust Case Control Consortium.

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Authors’ contributions
SE, WT, JW and AB conceived the study. EF and PM performed the genotyping, while AH and JB undertook the statistical analysis. YEAR and
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