The CYP27B1 variant associated with increased risk of autoimmune disease is underexpressed in tolerising dendritic cells

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ABSTRACT

Genome-wide association studies have identified a linkage disequilibrium block on chromosome 12 associated with multiple sclerosis (MS), type 1 diabetes and other autoimmune diseases. This block contains CYP27B1, which catalyses conversion of 25 Vitamin D3 to 1,25 Vitamin D3. Fine mapping analysis has failed to identify which of the 17 genes in this block is most associated with MS. We have previously used a functional approach to identify the causal gene. We showed that the expression of several genes in this block in whole blood is highly associated with the MS risk allele, but not CYP27B1. Here we show that CYP27B1 is predominantly expressed in dendritic cells (DCs). Its expression in these cells is necessary for their response to Vitamin D, which is known to upregulate pathways involved in generating a tolerogenic DC phenotype. Here we utilize a differentiation protocol to generate inflammatory (DC1) and tolerogenic (DC2) dendritic cells, and show that for the MS risk allele CYP27B1 is underexpressed in dendritic cells, especially DC2s. Of the other Chr12 LD block genes expressed in these cells, only METT21B expression was as affected by genotype. Another gene associated with autoimmune diseases, CYP24A1, catabolises 1,25 Vitamin D3, and is predominantly expressed in DCs, but equally between DC1s and DC2s. Overall, these data are consistent with the hypothesis that reduced vitamin D pathway gene upregulation in DC2s of carriers of the risk haplotype of CYP27B1 contributes to autoimmune diseases. These data support therapeutic approaches aimed at targeting Vitamin D effects on DCs.
INTRODUCTION

Up to 5% of the population of western countries develop autoimmune diseases (1). They result from an interaction between genetic and environmental factors. For the latter, a major advance has been confirmation that vitamin D (VitD) deficiency is associated with an increased risk of a range of autoimmune disorders (2). As further support, VitD supplementation has been shown to ameliorate disease in the animal models of autoimmune encephalomyelitis, collagen-induced arthritis, type 1 diabetes mellitus (T1D), inflammatory bowel disease, autoimmune thyroiditis, and systemic lupus erythematosus (SLE) (3).

The importance of VitD is now further implicated by genetic evidence. Large genome wide association studies (GWAS) have recently been completed with the statistical power to identify gene variants associated with autoimmune diseases. In multiple sclerosis (MS), 57 genetic regions were identified as associated with MS with genome wide significance (4). These included two genes regulating VitD activation, CYP27B1 and CYP24A1. CYP27B1 is in a linkage disequilibrium block of 17 genes on chromosome 12 (Chr12 LD block, Supp Fig. 1) originally identified as associated with MS in 2009 (5). Deep sequencing and fine mapping (6) and exon sequencing in affected kindreds (7) failed to establish which of the 17 genes was responsible. CYP24A1 is the only gene in the linkage block around SNP rs2248359, which is associated with MS. These findings provide the best genetic evidence to date that VitD has a causal association with MS. From GWAS, CYP27B1 and CYP24A1 are also associated with Type 1 diabetes (8, 9) and several other autoimmune diseases (10). Further, genes associated with autoimmune diseases (MS, T1D, rheumatoid arthritis, Crohn’s disease, SLE) have an over-representation of genomic VitD receptor (VDR) binding sites, as determined by ChIP-seq (11). VitD is used as a treatment by many with MS and other autoimmune diseases, and is currently being tested in clinical(12)
The main source of VitD comes from conversion of 7 dehydrocholesterol to pre-VitD in the skin by the action of UVB from sunlight, although VitD can also be obtained from the diet. For activation, it needs to be further converted by hydroxylation to 25OHD3 (25D3), usually in the liver, and finally to 1,25OHD3 (1,25D3), in certain cell subsets, by CYP27B1. 1,25D3 exerts its effects through the VDR, which acts as a ligand-activated transcription factor by binding to VitD response elements in the promoter regions of VitD responsive genes. The VDR is expressed in antigen-presenting cells and activated lymphocytes. CYP24A1 inactivates 1,25D3 and is itself a VDR target gene, thus providing a feedback loop, so that its VDR induction downregulates VDR function. These processes can occur in many cell types and impact on many cell processes, including immunoregulation (13). CYP27B1 is also expressed in cells in the brain where VDR-mediated pathways are postulated to contribute to a range of neurologic and neurophysiological disorders (14).

Although the evidence for VitD involvement in autoimmune pathogenesis is increasing, the genetic evidence is still equivocal because of the unresolved linkage disequilibrium block. The tight linkage disequilibrium of this block of genes and SNPs has confounded the identification of the gene driving the association with disease. We have hypothesised that the gene whose expression is most associated with the risk genotype is the functional one. In earlier work we had identified that expression of many genes of the Chr12 LD block in whole blood, which is predominantly lymphoid cells, are reduced for carriers of the rs10877012 risk allele, but not CYP27B1, which is scarcely expressed in these cells (15). The SNP rs108779012 is one of many that could be used to tag the risk allele (Supp Fig 1). In this study, we have now interrogated the expression of the LD block genes in myeloid cells. We have used ex vivo monocytes, plasmacytoid and myeloid DCs; and in vitro differentiation to model the inflammatory and tolerogenic DCs which control CD4 and CD8 differentiation, thought to be pivotal in MS pathogenesis.(16, 17) CYP27B1 expression was one of the most
genotype dependent, and the reduced expression of the risk allele of CYP27B1 in tolerising dendritic cells is consistent with reduced VitD function contributing to autoimmune disease risk. CYP24A1 was also predominantly expressed in DCs, and the balance of expression of these two genes likely contributes to regulation of DC phenotype and immune response.

RESULTS

Immune cell subsets important in MS susceptibility

The recent IMSGC MS GWAS (4) identified 57 SNPs associated with MS at genome-wide significant P values. To find the basis for the genetic association of any gene with MS a logical first step is to identify the cell subset in which it is predominantly expressed. As working hypotheses, we propose that the cell subset in which gene expression of the associated gene is most likely to be pathogenically relevant is that in which the genotype most affects expression and function; and that for genes from an LD block the gene most likely to be contributing to the association with susceptibility is the one for which genotype most affects expression. Taking the genes identified as most likely to be responsible for the association from (4), we interrogated their expression in 18 immune cell subsets and in oligodendrocytes, the brain cells destroyed in MS, and neuronal cells (Figure 1A). In all cases, expression was dominant in immune cells. Most genes were predominantly from a subset of immune cells, with genes such as THEMIS, CD6, MYC tagging T cells, CXCR5 tagging B cells, and many genes tagging myeloid cells. Many genes were expressed in multiple immune cell subsets. Overall, this suggests MS susceptibility is due to the interaction of all these immune cell subsets, but that genotype effects are likely to be mediated by the subset in which the gene is predominantly expressed.
Expression of Chr12 LD Block Genes in Immune Cell Subsets

For the genes of the Chr12 LD block, METT21B expression was by far the most associated with genotype in whole blood (15), where it is expressed at high levels in lymphocytes. Myeloid cells make up less than 10% of blood leukocytes. For these immune cells the association of genotype with gene expression is difficult to assess in blood, but these cells, particularly antigen presenting cells, of which DCs are the most potent, have long been considered to be involved in driving MS pathogenesis (16). From *in silico* data CYP27B1 and some other Chr12LD genes are most highly expressed in activated DCs in humans (18, 19). We confirmed this in immune cell subsets derived either directly from blood using Ficoll gradients, followed by purification with commercial magnetic bead kits, or culture in accepted conditions (Fig 1B, Fig 2). Of the examined immune cell subsets, we found the genes segregated into either those expressed predominantly in myeloid cells (CYP27B1, TSPAN31, METTL1, TSFM (also Th2), METTL21B, AVIL and OS9), differentiated T cells (CDK4, KIF5A, B4GALNT1, SLC26A10), or T cells and NK cells (AGAP2, ARHGEF5, CTDSP2, PIP4K2C, DTX3) (Figure 1B).

Dendritic Cell Differentiation

DCs are rare in blood, but common in difficult-to-sample sites such as lymph nodes and tissues. Fortunately, well established models for DC subsets exist, based on *in vitro* differentiation from CD14 monocytes, which are common in whole blood. We have used the method of Kalinsky (17), using interferon gamma (IFNG) to produce inflammatory DCs (DC1s), and, modified the method of Rani (20) to use interferon beta (IFNB), instead of TGFβ for generation of tolerogenic DCs (DC2s), since it produced a more polarised DC (greater difference between IL10 and IL12) (Fig 3, 4). Media and LPS did not alter cultured cell morphology, but 24hrs after addition of IFNG to culture, cells produced long dendrites...
(DC1); or after addition of IFNB, smaller dendrites (DC2s) (Fig 3). DC1s produced less IL10 than DC2s, at the mRNA and protein level (Fig 4) and the ratio of IL10 to IL12p40 mRNA was significantly higher in DC2s (Supp Fig 3). IL12p35 and IL12p40 were equally expressed in DC1 and DC2s, and at much higher levels than in CD14s (p). CYP27B1 and CYP24A1 genes were more highly expressed in DCs than CD14 cells, and CYP27B1 highest in DC2s (p)(Fig. 4). Notably, PBMCs treated with IFNB, or activated with PMA/ionomycin, do not upregulate CYP27B1 (Supp Fig 4), confirming the upregulation of CYP27B1 is a DC2 effect, not directly due to response to IFNB. Also, neither CYP27B1 nor CYP24A1 are recorded as interferon sensitive on the interferome(www.interferome.org). We also tested for a direct link between CYP27B1 expression and IL10, the key tolerogenic cytokine. They were strongly positively correlated (Pearson’s R of 0.5, p<0.0001; Fig. 5).

The effect of MS risk genotype on gene expression of Chr12 LD Block genes in DCs

CD14 cells were isolated, and DC1 and DC2 cells differentiated from 49 healthy controls of known rs10877012 genotype. Of the Chr12 LD block myeloid-expressed genes, CYP27B1 was the most highly expressed, especially in DC2s. METT1, METT21B, MARCH9 and TSFM were also more highly expressed in DC2s than DC1s (p<0.001). TSPAN31 was most highly expressed in CD14s (p<0.0001)(paired t tests, two tailed, Fig 6A). Expression of CYP27B1 and METT21B were significantly increased for the protective genotype, rs10877012 T, in DC2s (Fig 6B, 6C). From the cell subset RNAseq data, the splice isoforms present for each gene were inferred. In DCs, only one isoform was seen for CYP27B1, CYP24A1 and 8 other genes. Multiple isoforms were seen for METT21B and TSFM in DC2s (Fig 7, Supp Fig 2). We then tested if genotype affected the isoform usage for METT21B and TSFM in DC2s. For the former it did (p<0.0047, t test, two tailed) and a trend was seen for TSFM.
The effect of MS risk genotype on CYP24A1 gene expression

CYP27B1 activates 25VitD3, CYP24A1 catabolises it, and is also associated with MS. We tested which immune cells expressed CYP24A1, and if genotype of the MS-associated SNP, rs2248359 affected expression. Like CYP27B1, it was also more highly expressed in myeloid cells than lymphoid cells, and of them most highly expressed in DCs, but not higher in DC2s than DC1s (Fig 8A). Genotype did not affect expression (Fig 8B).

DISCUSSION

Over 50 gene regions have now been identified which affect MS susceptibility. Yet in most cases the associated SNPs tag LD blocks which compromise further studies to identify the functional significance of these genetic associations. One approach is to focus on genes from the LD block that are expressed in pathogenically relevant cells, and test the effect of the risk allele on expression. In this study we have tested the expression of genes from the chromosome 12 rs10877012 LD block in myeloid cell subsets, and demonstrated that for several of these genes expression is highest in DCs, where the protective haplotype is generally more highly expressed. This is especially so for CYP27B1. CYP27B1’s partner in controlling VitD levels in cells, CYP24A1, is both associated with MS and also predominantly expressed in DCs. CYP27B1 expression is higher in the tolerogenic DCs, DC2s, and CYP24A1, which has an opposite effect on cellular 1,25D3 levels, is not.

Regulation of expression of this linkage block in multiple cell types may be due to the same regulatory region responding to cell subset specific transcription factor complexes, chromatin remodelling and DNA methylation enzymes. We have shown here that the splicing of genes in the LD block was genotype specific for METT21B in DC2s, and so mRNA isoform usage
for this, and potentially the other genes in the LD block, may contribute to MS risk. Only one isoform of CYP27B1 is described in the RefSeq database (build 57) at this time. Alcina et al (21) have recently reviewed transcriptional control across this region from the ENCODE project, published genome-wide eQTL studies from different immune cell types, and they used luciferase constructs in B cells for different candidate SNPs from the risk haplotype to identify functional SNPs. From the former there were strong heritable effects on gene expression in the LD block, with the direction and extent of association dependent on the cell type assayed and gene studied. From the latter, SNP rs10807712 had the largest effect on transcription. Notably though, many other SNPs from this haplotype block have not been tested, and the genotype association with transcription might prove to be cell context specific. We have previously reported that in blood expression of the gene METT21B (methyltransferase protein 21B, previously known as FAM119B) is by far the most highly associated with the risk haplotype. The major alleles of SNPs rs10877012 (G) and rs703842 (A) tag the CYP27B1 haplotype associated with increased risk of autoimmune disease (4), and decreased expression in whole blood and DCs. Overall, on the basis of association of genotype with gene expression alone, the strongest candidates for driving the association of the Chr12 LD block haplotype with autoimmune diseases are CYP27B1 and METT21B.

Of the two, there is strong evidence from other approaches to support CYP27B1. It activates VitD, deficiency of which is associated with an increased risk of a range of autoimmune disorders (2). VitD supplementation has been shown to ameliorate disease in the animal models of autoimmune encephalomyelitis, collagen-induced arthritis, T1D, inflammatory bowel disease, autoimmune thyroiditis, and SLE (3). The gene which inactivates 1,25D3, CYP24A1, is also associated with MS, and of immune cells it is also produced predominantly in DCs, though less in tolerogenic DCs than CYP27B1. The immunomodulatory effects of
VitD3 are mediated mainly through its action on dendritic cells, where these effects are most potent and well described (22, 23). The VDR pathway activates genes which induce DCs with a tolerogenic phenotype (24, 25), reduce autoimmune disease in mouse models (26), and 1,25D3 can be used to produce clinical grade tolerogenic DCs (27). Other immune cells have limited CYP27B1 expression, but they could potentially respond to an exogenous source of 1,25D3 and suppress autoimmunity by inducing regulatory T cells (Tregs) (28) and natural killer T cells (29), by inhibiting pro-inflammatory Th17 cells (30), or by inducing apoptosis of activated T cells (31, 32). The production of anti-inflammatory cytokines from non-immune cells as well as immune cells may also be affected by VitD therapy (3). The variant of CYP27B1 that increases risk of autoimmune disease, is produced at lower levels in DC2s than the protective variant, which is consistent with less VDR gene activation and tolerogenic function in these cells. Notably CYP27B1 mRNA levels correlated with IL10 mRNA levels in DC2s. This suggests they are on shared regulatory pathways. This higher IL10 production may be a consequence of the higher production of 1,25D3 with higher CYP27B1 expression, and subsequent upregulation of tolerogenic genes mediated by the vitamin D receptor. This is consistent with a tolerogenic effect of the more highly expressed protective haplotype.

The circulating level of 25D3, the best measure of VitD stores, is under genetic influence. In a very large GWAS (n=33,300), four genes affecting 25D3 levels in Europeans were identified (33): GC, DHCR7, CYP2R1 and CYP24A1. The first three have not been associated with autoimmune disease at genome-wide significance and the CYP24A1 SNP associated with 25D3 levels (rs6013897) is different from, and not in LD with, the CYP24A1 MS risk SNP (rs2248359). These data suggest that the genetic control of serum level of 25D3 is not as important to MS susceptibility as the intra-cellular regulation of 1,25D3; and that supplements, which are used to increase serum 25D3 levels, may not be targeting the cellular
functions relevant to pathogenesis.

An array of drugs has already been designed by Pharma to manipulate the VitD3 pathway (13). VDR agonist ligands carry a risk of hypercalcaemia, and loss of activity due to VDR-mediated up-regulation of CYP24A1 leading to ligand catabolism. VDR ligands that minimise these problems are under development (13, 34). Targeting the VDR agonist to the cell type relevant to the therapeutic effect is a logical approach to therapy, but this necessitates detailed mechanistic knowledge of the disease process. To date, many clinical trials have been undertaken in MS, but only with the natural ligand (1,25D3) or its immediate precursor (25D3), with mixed results (35-37).

These data indicate that the basis for the genetic association of variants of CYP27B1 and CYP24A1 with MS are likely due to their expression in dendritic cells altering the balance of tolerogenic and inflammatory DCs. This supports the concept that therapeutic VitD should target DCs, and that *ex vivo* manipulation of DCs with VitD for autologous transplantation might be suitable as a treatment for MS and other autoimmune diseases affected by the same risk variants of these genes.
MATERIALS AND METHODS

Study Subjects
Approximately 25 mL of peripheral blood was collected from 49 healthy control subjects between the ages of 25-68. Subjects were laboratory personnel not on any immunomodulatory medications. The Human Research Ethics Committee from the Sydney West Area Health Service approved this study. Written informed consent was given by all participants.

RNA Analysis of Immune, Oligodendrocyte, and Neuronal Cell Subsets
Next generation mRNA sequencing (RNA-seq) was performed for the following immune cell subsets: peripheral blood mononuclear cells (PBMCs) and neutrophils ex Ficoll; then CD4, CD4+CD45RA+, CD4+CD45RO+, CD8+ T cells, and Tregs purified ex vivo by magnetic beads from the PBMC fraction; Th1, Th2, Th17 cultured in vitro, CD19+ B cells, Natural Killer cells (NK: CD3-CD56+), plasmacytoid DCs (pDC: CD303+CD304+CD123+CD11c-), myeloid DCs (mDC; CD19-CD1chi), and CD14 monocytes were purified by magnetic beads, then cells derived from CD14 cells in vitro: monocyte derived DCs (MDDC, IL4/GMCSF), monocyte derived macrophages (MDM, IL4/M-CSF), followed by LPS for 48 hours, producing DCs (CD14-HLA- DRhiCD11c+CD40hiCD80+). If treated at day 6 with IFNG (1000u/ml) these become DC1s (IL12hi), or with IFNB, DC2s (IL10hi). RNA-seq was also performed for the following non immune cells; 4-β-phorbol 12-myristate 13-acetate (PMA)-differentiated human MO3.13 cell line (kind gift from Dr Gilles Guillemin) used as a model of myelin basic protein-expressing human oligodendrocytes; retinoic acid-differentiated human SH-SY5Y cells (www.ATCC.org) used as a model of dopaminergic neuronal cells (38). RNA was isolated from these cell subsets using Qiagen RNeasy Mini kit and RNA was prepared for sequencing on Illumina HiSeq 2000 using the Illumina TruSeq RNA sample
preparation kit V1. Raw sequence data was aligned to the UCSC human reference genome (hg19) using the Tophat software package (39). Aligned sequencing reads were summarised to both counts per gene and counts per transcript (as per RefSeq Release 57) using the RAEM procedure (40) and RPKM values calculated in SAMMate (41) (v 2.6.1). RPKM values were transformed by quantile normalisation prior to visualisation. Visualisation was performed by performing hierarchical clustering using one minus correlation and average linkage metrics (BRB ArrayTools v 4.2.1(42)).

Myeloid Cell Culture

Peripheral blood mononucleocytes (PBMCs) were isolated from EDTA-treated whole blood of healthy controls using Ficoll Paque Plus (GE Healthcare, Sweden). CD14+ cells were purified from the PBMCs using the Human CD14 Positive Selection Kit (Stem Cell Technologies, Canada) according to the manufacturer’s instructions. Cells were plated at a density of $2 \times 10^5$ cells per well in X-Vivo15 medium (Lonza, Switzerland) with 2% AB serum in four sets of triplicate 96-well flat-bottom plate wells and treated with three changes of GMCSF (70 ng per ml) and IL-4 (10 ng per ml) over six days. On day 6, cells were treated with medium only, LPS (250 ng per ml) only, LPS with IFN-γ (10 ng per ml) or LPS with IFN-β (1000 IU per ml). After a further 48 hours, cells from single wells of each treatment were removed to Cells-to-Signal Lysis Buffer (Ambion) for purification of RNA, while cells from the remaining two wells were removed for flow cytometry analysis.

RTPCR

RNA was purified from samples using an RNeasy Minikit (Qiagen) and reverse transcribed using QScript reverse transcriptase (Quanta Biosciences), each according to manufacturer’s instructions. cDNA was diluted ten-fold with RNase-free/DNase-free water and analysed by
real time PCR using POWER Sybr Green Mastermix (Applied Biosystems) and primer sets for GAPDH, IDO, IL-10, IL-12 p35, IL-12 p40, CYP27B1 and CYP24A1:

GAPDH: Fwd 5’-TCCACCACCCTGTTGCTGTA-3’, Rev 5’-ACCACAGTCCATGCCATCAC-3’;
IL-10: Fwd 5’-TTACCTGGAGGAGGTGATGC-3’, Rev 5’-GGCCTTGCTCTTCTTTTCAC-3’;
IL-12p35: Fwd 5’-CCTCCTGGACCACCTCAGTTTG-3’, Rev 5’-CCTCCTGGACCACCTCAGTTTG-3’;
IL-12p40: Fwd 5’-TCAAAGAGTTTGGAGATGCTGGC-3’, Rev 5’-TGATGATGTCCCTGATGAAGAAGC-3’;
CYP27B1: Fwd 5’-TTTGCATCTCTTCCTTTTG-3’, Rev 5’-AGGTTGATGCTCCTTTCAGG-3’;
CYP24A1: Fwd 5’-CATCATGGCCATCAAAACAAT-3’, Rev 5’-GCAGCTCGACTGGAGTGAC-3’.
The probes used to quantify the expression of Chr12LD block genes expressed in myeloid cells: METTL21B, METTL1, MARCH9, TSPAN31 and TSFM, and the methods used for quantification of METT21B and TSFM isoforms are described in the supplementary information.

Genotyping

Genomic DNA was isolated from 200uL peripheral blood using QIAmp DNA Blood Mini Kit (Qiagen). DNA was diluted ten-fold with RNase-free/DNase-free water and analysed by real time PCR using Taqman Genotyping Master Mix (Applied Biosystems) and primer sets for CYP27B1 (rs10877012) and CYP24A1 (rs2248359).

Statistics

Comparisons were made using Mann Whitney test or paired, two tailed T tests between carrier and non-carrier genotypes, and was considered significant if p≤0.05, as described in the figure legends. All real time PCR RNA expression was measured relative to GAPDH and, when applicable, error margins were calculated using standard error.
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CONFLICTS OF INTEREST

The authors declare no conflict of interests.

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LEGENDS TO FIGURES

Figure 1. A. Expression profile of genes associated with multiple sclerosis from the IMMSGC GWAS (4). Cell subsets were isolated and mRNA quantified by RNA-seq as described in Methods and Supplementary Methods. Columns are cell subsets, rows are genes. High expression is orange, low expression is blue. B. Expression profile of genes from the Chromosome 12 rs10877012 linkage disequilibrium block in the same cell subsets.

Figure 2. CYP27B1 expression in Immune Cell Subsets. Cell subset derivation is described in Methods and Supplementary Methods. Expression is measured relative to GAPDH (n=3/cell subset, including at least 1 heterozygote rs10877012; n=2 for plasmacytoid dendritic cells).

Figure 3. Validation of DC1 and DC2 phenotypes by morphological criteria. CD14 cells treated with IL4 and GM-CSF for 6 days then with A. media alone or B. LPS, showed small to no dendrites signifying weak or no antigen presentation. Cells treated from day 6 with either C. LPS+IFNG (DC1) or D. LPS+IFNB (DC2) show long dendrites, signifying stronger antigen presentation with the most pronounced dendrites on LPS+IFNG treated cultures. Pictures taken at 200X magnification on a Leica upright fluorescence microscope.

Figure 4. Validation of DC1 and DC2 by cytokine mRNA production. A. DC1s produce less IL10 than DC2s (n=49, p<0.001). B. IL10 and IL12p70 proteins in media of undifferentiated DCs (media), DC1 (IFNG) and DC2 (IFNB) cells. Typical example shown. C. CYP27B1 is more highly expressed in DCs than CD14s, and more highly expressed in DC2 than DC1s (n=49, p<0.00001, both comparisons, paired T test). D. CYP24A1 was not detectable in
CD14s, and not different between DCs. (n=49, but in 11 no signal detected (no genotype bias in undetected)). Measurements relative to GAPDH. Paired t tests used. Standard error bars shown.

**Figure 5.** CYP27B1 mRNA expression was correlated with IL10 mRNA expression in DC2 cells. Correlation was tested using Pearson’s linear correlation, comparing –log10 (2^th) values (R=0.50, p<0.0001).

**Figure 6.** A. Comparative expression of Chr12 rs10877012 LD block genes in CD14 monocytes and DC1 and DC2s (n>=40 for each gene). B. Effect of SNP rs10877012 genotype on expression of Chr12 rs10877012 LD block genes in DC2s (n=49 for all, -log_{10}(p value), Mann Whitney test, used to compare association of TT/GTvsGG for the different genes). C. Genotype of CYP27B1 SNP rs10877012 appears to be associated with expression in DC2s (n=49, TT/GTvsGG, p<0.012, Mann Whitney test), and DC1s (p<0.03), with lower expression of the variant (G) increasing susceptibility to MS.

**Figure 7.** A. Splice isoforms of METT21B in immune cell subsets. Refseq isoform codes in the legends. B. Genotype association of the proportion of the two METT21B isoforms in DC2s (p=0.0047, T test (two tailed) of TT vs GG/GT). C. Splice isoforms of TSFM in immune cell subsets. D. Genotype association of the proportion of the two TSFM isoforms in DC2s (ns).

**Figure 8.** A. CYP24A1 expression in immune cell subsets largely mirrors that of CYP27B1 (n>=3/subset) B. CYP24A1 SNP rs2248359 genotype appears not to affect expression in DCs (n=49 examined, 40 (DC1) and 41 (DC2) plotted in figure as no signal detected in 9 and 8 respectively (no genotype bias in these 9 and 8). p value from Mann Whitney test.
ABBREVIATIONS

VitD: Vitamin D

VDR: Vitamin D Receptor

DC; dendritic cell

CHR12LD Block chromosome: 12 linkage disequilibrium block

MS: multiple sclerosis

T1D: type 1 diabetes

SLE: systemic lupus erythematosus

Gene names are abbreviated according to the Hugo Gene Nomenclature Committee