#### 1 Whole Genome Sequencing of Primary Immunodeficiency reveals a role for common and rare

### 2 variants in coding and non-coding sequences

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#### 59 Abstract

60

61 Primary immunodeficiency (PID) is characterised by recurrent and often life-threatening infections,

62 autoimmunity and cancer, and it presents major diagnostic and therapeutic challenges. Although the

- 63 most severe forms present in early childhood, the majority of patients present in adulthood, typically
- 64 with no apparent family history and a variable clinical phenotype of widespread immune dysregulation:
- about 25% of patients have autoimmune disease, allergy is prevalent, and up to 10% develop lymphoid
- 66 malignancies. Consequently, in sporadic PID genetic diagnosis is difficult and the role of genetics is not
- 67 well defined. We addressed these challenges by performing whole genome sequencing (WGS) of a large
- 68 PID cohort of 1,318 subjects. Analysis of coding regions of 886 index cases found disease-causing
- 69 mutations in known monogenic PID genes in 8.2%, while a Bayesian approach (BeviMed<sup>1</sup>) identified
- 70 multiple potential new disease-associated genes. Exploration of the non-coding space revealed
- 71 deletions in regulatory regions which contribute to disease causation. Finally, a genome-wide
- 72 association study (GWAS) identified novel PID-associated loci and uncovered evidence for co-localisation
- of, and interplay between, novel high penetrance monogenic variants and common variants (at the
- 74 *PTPN2* and *SOCS1* loci). This begins to explain the contribution of common variants to variable
- penetrance and phenotypic complexity in PID. Thus, a cohort-based WGS approach to PID diagnosis can
- 76 increase diagnostic yield while deepening our understanding of the key pathways determining variation
- 77 in human immune responsiveness.

78

- 79 The phenotypic heterogeneity of PID leads to diagnostic difficulty, and almost certainly to an
- 80 underestimation of its true incidence. Our cohort reflects this heterogeneity, though it is dominated by
- 81 adult onset, sporadic antibody deficiency associated PID (AD-PID: comprising Common Variable
- Immunodeficiency (CVID), Combined Immunodeficiency (CID) and isolated antibody deficiency). 82
- 83 Identifying a specific genetic cause of PID can facilitate definitive treatment including haematopoietic
- stem cell transplantation, genetic counselling, and the possibility of gene-specific therapy<sup>2–4</sup> while 84
- 85 contributing to our understanding of the human immune system<sup>5</sup>. Unfortunately, only 29% of patients
- with PID receive a genetic diagnosis<sup>6</sup>. The lowest diagnosis rate is in patients who present as adults, 86
- 87 have no apparent family history, and in whom matching the clinical phenotype to a known genetic cause
- 88 is difficult, as the latter can be surprisingly variable even in patients with the same genetic defect (in the
- 89 UK PID cohort 78% of cases are adult and 76% sporadic<sup>6</sup>). Moreover, while over 300 monogenic causes 90 of PID have been described<sup>7</sup>, the genotype-phenotype correlation in PID is complex. In CVID, for
- 91 example, pathogenic variants in TACI (TNFRSF13B) occur in 10% of patients but typically have low
- 92 disease penetration, appearing to act as disease modifiers<sup>8</sup>. Furthermore, a common variant analysis of
- 93 CVID identified two disease-associated loci, raising the possibility that common variants may impact
- 94 upon clinical presentation<sup>9</sup>. We therefore investigated whether applying WGS across a "real world" PID
- 95 cohort might illuminate the complex genetics of the range of conditions collectively termed PID.
- 96

#### 97 **Patient cohort**

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99 974 sporadic and familial PID patients, and 344 unaffected relatives, were recruited by collaborators as 100 part of the United Kingdom NIHR BioResource - Rare Diseases program (NBR-RD; Supplementary Note). 101 Of these, 886 were index cases who fell into one of the diagnostic categories of the European Society for 102 Immunodeficiencies (ESID) registry diagnostic criteria (Fig. 1a; Supplementary Table 1). This cohort represents a third of CVID and half of CID patients registered in the UK<sup>10</sup>. Paediatric and familial cases 103 104 were less frequent, in part reflecting prior genetic testing of more severe cases (Supplementary Fig. 1). Clinical phenotypes were dominated by adult-onset sporadic AD-PID: all had recurrent infections, 28% 105 106 had autoimmunity, and 8% had malignancy (Fig. 1a-b, Supplementary Table 2), mirroring the UK

- national PID registry<sup>6</sup>. 107
- 108

#### 109 Identification of Pathogenic Variants in Known Genes

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We analysed coding regions of genes previously causally associated with PID<sup>11</sup> (Methods). We identified 111 112 85 potentially causal variants in 73 index cases (8.2%) across 39 genes implicated in monogenic disease 113 (Fig. 1c; Supplementary Table 3). 60 patients (6.8%) had a previously reported pathogenic variant in the 114 disease modifier TACI (TNFRSF13B), increasing the diagnostic yield to 15.0% (133 patients). Interestingly, 115 5 patients with a monogenic diagnosis (in BTK, LRBA, MAGT1, RAG2, SMARCAL1) also had a pathogenic 116 TACI variant. The diagnostic yield rose to 17.0% (151 patients) once novel causal variants in NFKB1 and 117 ARPC1B, associated with PID only after our initial analysis, were included. Of the 85 monogenic variants 118 we reported, 51 (60%) had not been previously described (Supplementary Table 3), and 4 were structural variants, including a single exon deletion, unlikely to have been detected by whole exome 119 120 sequencing<sup>12</sup>.

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- 122 We observed divergence from an expected clinical phenotype for causal variants in 14 genes: for
- instance, only 4 of the 8 STAT1 patients had the pathognomonic chronic mucocutaneous candidiasis<sup>13,14</sup>. 123
- 124 A more remarkable example of phenotypic complexity was the case of a 40 year-old patient presenting
- 125 with specific antibody deficiency and a premature stop variant at Arg328 in X-linked IL2RG, a defect
- 126 expected to cause absent T and NK cells and death in infancy. We found that the mild phenotype could

be ascribed to several independent somatic changes that reversed the premature stop codon, restoring
both T and NK cell lineages (Fig. 1d and Supplementary Fig. 2).

129

130 Since many PID-associated genes were initially discovered in a small number of typically familial cases, it

is perhaps not surprising that the phenotypes described do not reflect true clinical diversity. Thus, a

132 cohort-based WGS approach to PID can provide a significant diagnostic yield even in a predominantly

133 pre-screened and sporadic cohort, allows diagnoses which are not constrained by pre-existing

assumptions about genotype-phenotype relationships, and suggests caution in the use of clinical

135 phenotype in targeted gene screening and when interpreting PID genetic data.

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## 137 An approach to identifying new PID-associated genes in a WGS cohort

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139 We next sought to determine whether the cohort-based WGS approach could identify new genetic

associations with PID. We developed a Bayesian inference procedure, named BeviMed<sup>1</sup>, to determine

141 posterior probabilities of association (PPA) between each gene and case/control status of the 886 index

142 cases and 9,283 unrelated controls (**Methods**). For each gene, we analysed variants with gnomAD minor

- allele frequency (MAF) <0.001 and Combined Annotation Dependent Depletion (CADD) score >=10.
- 144 Genes with PPA>=0.18 are shown in **Fig. 1e**. There was a strong enrichment for known PID genes

145 (Wilcoxon P<1x10<sup>-200</sup>), supporting this statistical approach. Two novel BeviMed-identified genes were

subsequently causally associated with PID. *NFKB1* had the strongest probability of disease association

(PPA=1.0), driven by truncating heterozygous variants in 13 patients. Subsequent assessment of co segregation, protein expression, and B cell phenotype in pedigrees established these as disease-causing

- variants, and consequently loss of function variants in *NFKB1* as the most common monogenic cause of
- 150 CVID<sup>15</sup>. Evidence of association of *ARPC1B* with PID (PPA=0.18) was driven by 2 functionally validated
- 151 cases, one homozygous for a complex InDel<sup>16</sup> and the other described below.

The discovery of both known and subsequently validated new PID genes using BeviMed underlines its
 effectiveness in cohorts of unrelated patients with sporadic disease. Many candidate genes identified by
 BeviMed remain to be functionally validated and, as the PID cohort grows, even very rare causes of PID
 (e.g. affecting 0.2% of cases) will be detectable with a high positive predictive value (Supplementary Fig.
 3).

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# 158 Identification of regulatory elements contributing to PID

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Sequence variation within non-coding regions of the genome can have profound effects on spatial and 160 temporal gene expression<sup>17</sup> and would be expected to contribute to PID susceptibility. We combined 161 rare variant and deletion events with a tissue-relevant catalogue of cis-regulatory elements (CREs)<sup>18</sup> 162 generated using promoter capture Hi-C (pcHi-C)<sup>19</sup> in matching tissues to prioritise putative causal PID 163 genes (Fig. 2a). Being underpowered to detect single nucleotide variants affecting CREs<sup>20</sup>, we limited our 164 initial analysis to rare structural variants (SV) overlapping exon, promoter or 'super-enhancer' CREs of 165 166 known PID genes. No homozygous deletion events affecting CREs were identified, so we sought CRE SV 167 deletions that might cause disease through a candidate compound heterozygote (cHET) mechanism with 168 either a heterozygous rare coding variant or another SV in a pcHi-C linked gene (Fig. 2a). Out of 22,296 169 candidate cHET deletion events, after filtering by MAF, functional score and known PID gene status, we obtained 10 events; the functional follow-up of three is described (Fig. 2b). 170 171

172 The *LRBA* and *DOCK8* cHET variants (**Supplementary Fig. 4**) were functionally validated; the former was 173 demonstrated to result in impaired surface CTLA-4 expression on Treg cells (**Supplementary Fig. 5**) whilst the latter led to DOCK8 deficiency as confirmed by flow cytometry (data not shown). Although in
these two cases SV deletions encompassed both non-coding CREs and coding exons, the use of WGS PID
cohorts to detect a contribution of CREs confined to the non-coding space would represent a major

- advance in PID pathogenesis and diagnosis. *ARPC1B* fulfilled this criterion, with its BeviMed association
- partially driven by a patient cHET for a novel p.Leu247Glyfs\*25 variant resulting in a premature stop,
- and a 9Kb deletion spanning the promoter region including an untranslated first exon (**Fig. 2c**) that has
- 180 no coverage in the ExAC database (http://exac.broadinstitute.org). Two first-degree relatives were
- 181 heterozygous for the frameshift variant, and two for the promoter deletion (**Fig. 2d**). Western blotting
- demonstrated complete absence of ARPC1B (**Fig. 2e**) and, consistent with previous reports<sup>21</sup>, raised
- 183 ARPC1A in platelets. *ARPC1B* mRNA was almost absent from mononuclear cells in the cHET patient and
- reduced in a clinically unaffected sister carrying the frameshift mutation (Fig. 2f). An allele specific
   expression assay demonstrated that the promoter deletion essentially abolished mRNA expression (Fig.
- 186 **2g,h**).

187 These examples show the utility of WGS for detecting compound heterozygosity for a coding variant and

a non-coding CRE deletion, and demonstrate a further advantage of a WGS approach to PID diagnosis.

189 Improvements in analysis methodology, cohort size and better annotation of regulatory regions will be

190 required to explore the non-coding space more fully and discover new disease-causing genetic variants.

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# 192 WGS identifies PID-associated telomere shortening

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194 A striking example of WGS data providing more than just the linear genomic sequence is telomere

195 length estimation from mapped and unmapped reads<sup>22</sup>. We validated this method by showing

196 correlation with gender (Fig. 3a) and a particularly strong correlation with age (Supplementary Fig. 6) in

197 3,313 NBR-RD subjects (**Methods**). We demonstrated the effectiveness of this, the first large-scale

application of WGS-based telomere length estimation, by replicating an association with the telomerase

199 RNA component gene (*TERC*: Supplementary Table 4)<sup>23</sup> and identifying several PID cases with short
 200 telomeres (Fig. 3b). Given that disruption of telomerase genes can cause PID<sup>24</sup>, we looked for potentially

201 damaging coding variants in known telomere deficiency genes<sup>25</sup> in these PID cases, identifying 3 subjects

with novel variants potentially causative for telomerase deficiency (**Fig. 3b**). One had a homozygous

203 defect in telomerase reverse transcriptase (*TERT*), a subunit of the telomerase complex. Two male

siblings were found to have a hemizygous variant in dyskerin (*DKC1*), known to be associated with PID

and X-linked dyskeratosis congenita<sup>26</sup> (Fig. 3c). Therefore, WGS telomere length estimation can be used
 as an effective approach to identify PID patients with novel variants causing telomere shortening.

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# 208 GWAS of the WGS cohort reveals novel PID-associated loci

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The diverse clinical phenotype and variable within-family disease penetrance of PID may be in part due to stochastic events (e.g. unpredictable pathogen transmission) but may also have a genetic basis. We

therefore performed a GWAS of common SNPs (MAF>0.05), restricted to 733 AD-PID cases (Fig. 1a) to

reduce phenotypic heterogeneity, and 9,225 unrelated NBR-RD controls. We confirmed the known MHC

association and identified additional loci with suggestive association (Fig. 4a, Supplementary Fig. 7). A

GWAS of SNPs of intermediate frequency (0.005<MAF<0.05) identified a single locus incorporating

216 TNFRSF13B (Fig. 4a, Supplementary Table 5, Extended Data Fig. 1), for which the lead p.Cys104Arg

217 variant has been previously reported<sup>27</sup>.

218 To increase power, we conducted a fixed effect meta-analysis of the AD-PID GWAS with summary

statistics data from an ImmunoChip study of 778 CVID cases and 10,999 controls<sup>9</sup> (Fig. 4a,

220 **Supplementary Table 5**). This amplified the MHC and 16p13.13 associations<sup>9</sup>, found an additional locus

at 3p24.1 within the promoter region of *EOMES* (Extended Data Fig. 2), and a suggestive association at

222 18p11.21 proximal to *PTPN2* (Extended Data Fig. 3). Conditional analysis of the MHC locus revealed

independent signals at the Class I and Class II regions (**Supplementary Fig. 8**), driven by classical alleles

HLA-B\*08:01 and HLA-DRB1\*15:01 (Methods) with amino-acid changes known to impact upon peptide binding (Fig. 4b).

226 We next sought to examine, genome-wide, the enrichment of non-MHC AD-PID associations in 9 other diseases (Extended Data Table 1). We found significant enrichment for allergic (e.g. asthma) and 227 228 immune-mediated diseases (e.g. Crohn's disease), which was not evident in Type 2 diabetes or coronary 229 artery disease (Fig. 4c). This suggests that the common variant association between PID and other 230 immune-mediated diseases extends beyond the 4 genome-wide loci to multiple sub-genome-wide 231 associations, and that dysregulation of common pathways contributes to susceptibility to both. 232 Understanding the impact of these interrelationships will be a complex process. For example, while variants in the HLA-DRB1 and 16p13.13 loci increase the risk of both PID and autoimmunity, those at the 233 234 EOMES locus predispose to PID but protect from rheumatoid arthritis<sup>28</sup> (Extended Data Fig. 2).

235

236 Given this observed enrichment, we sought to investigate whether candidate genes identified through 237 large cohort association analysis of immune-mediated disease might have utility in prioritising novel 238 candidate genes harbouring rare coding variation causal for PID. We used the data-driven capture-HiC omnibus gene score (COGS) approach<sup>19</sup> to prioritise putative causal genes across the 4 non-MHC AD-PID 239 240 loci identified by our meta-analysis, and assessed across 11 immune-mediated diseases (Supplementary 241 Tables 5 and 6). Hypothesising that causal PID genes would be intolerant to protein-truncating variation, we computed an overall prioritisation score by taking the product of pLI (a measure of tolerance to loss 242 243 of gene function) and COGS gene scores for each disease. Six protein coding genes had an above 244 average prioritisation score in one or more diseases (Fig. 4d) which we examined for rare, potentially 245 causative variants within our cohort. We identified a single protein truncating variant in ETS1, SOCS1 246 and PTPN2 genes, all occurring exclusively in PID patients in the NBR-RD cohort. None of the genes are 247 recognised causes of PID despite their involvement in immune processes (Supplementary Discussion). 248 The two cases with SOCS1 and PTPN2 variants were analysed further.

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250 The patient with a heterozygous protein-truncating SOCS1 variant (p.Met161Alafs\*46) presented with 251 CVID complicated by lung and liver inflammation and B cell lymphopenia (Supplementary Discussion, 252 Supplementary Fig. 9). SOCS1 limits phosphorylation of targets including STAT1, and is a key regulator of IFN-y signalling. SOCS1 haploinsufficiency in mice leads to B lymphopenia<sup>29,30</sup>, immune-mediated liver 253 inflammation<sup>31</sup> and colitis<sup>32</sup>. In patient T cell blasts SOCS1 was deficient and IFN-y induced STAT1 254 255 phosphorylation was abnormal (Fig. 4e), consistent with SOCS1 haploinsufficiency causing PID. The 256 patient also carries the SOCS1 pcHiC-linked 16p13.13 risk-allele identified in the AD-PID GWAS 257 (Extended Data Fig. 4). Long read sequencing using Oxford Nanopore technology showed this to be in 258 trans with the novel SOCS1-truncating variant (Methods); such compound heterozygosity raises the 259 possibility that common and rare variants may combine to cause disease.

260

A more detailed example of an interplay between rare and common variants is provided by a family containing a novel *PTPN2* premature stop-gain at p.Glu291 and a common autoimmunity-associated variant (**Fig. 4f**). *PTPN2* encodes the non-receptor T-cell protein tyrosine phosphatase (TC-PTP) protein, that negatively regulates immune responses by dephosphorylation of the proteins mediating cytokine

signalling. *PTPN2* deficient mice are B cell lymphopenic<sup>33,34</sup>, while inducible haematopoietic deletion of

266 *PTPN2* leads to B and T cell proliferation and autoimmunity<sup>35</sup>. The novel truncating variant was

- identified in a "sporadic" index case presenting with CVID at age 20; he had B lymphopenia
- 268 (Supplementary Fig. 9), low IgG, symmetrical rheumatoid-like polyarthropathy, severe recurrent

269 bacterial infections, splenomegaly and inflammatory lung disease. His mother, also heterozygous for the 270 PTPN2 truncating variant, had systemic lupus erythematosus (SLE), insulin-dependent diabetes mellitus 271 diagnosed at 42, hypothyroidism and autoimmune neutropenia (Supplementary Discussion). Gain-offunction variants in STAT1 can present as CVID (Supplementary Table 3) and TC-PTP, like SOCS1, 272 273 reduces phosphorylated-STAT1 (Fig. 4g). Both mother and son demonstrated reduced TC-PTP expression and STAT1 hyperphosphorylation in T cell blasts, similar to the SOCS1 haploinsufficient patient above 274 275 and to known STAT1 GOF patients; abnormalities that were more pronounced in the PTPN2 index case 276 (Fig. 4h).

277 278 The index case, but not his mother, carried the G allele of variant rs2847297 at the PTPN2 locus, an expression quantitative trait locus (eQTL)<sup>36</sup> previously associated with rheumatoid arthritis<sup>37</sup>. His 279 280 brother, generally healthy apart from severe allergic nasal polyposis, was heterozygous at rs2847297 281 and did not inherit the rare variant (Fig. 4f). Allele-specific expression analysis demonstrated reduced 282 PTPN2 transcription from the rs2847297-G allele, explaining the lower expression of TC-PTP and greater 283 persistence of pSTAT1 in the index case compared to his mother (Fig. 4i). This in turn could explain the 284 variable disease penetrance in this family, with *PTPN2* haploinsufficiency alone driving autoimmunity in 285 the mother, but with the additional impact of the common variant on the index case causing immunodeficiency (and perhaps reducing the autoimmune phenotype). The family illustrates the power 286 287 of cohort-wide WGS approach to PID diagnosis, by revealing both a new monogenic cause of disease, 288 and how the interplay between common and rare genetic variants may contribute to the variable clinical 289 phenotypes of PID.

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291 In summary, we show that cohort-based WGS in PID is a powerful approach to provide immediate 292 diagnosis of known genetic defects, and to discover new coding and non-coding variants associated with 293 disease. Intriguingly, even with a limited sample size, we could explore the interface between common 294 and rare variant genetics, explaining why PID encompasses such a complex range of clinical syndromes 295 of variable penetrance. Increasing cohort size will be crucial for powering the analyses needed to 296 identify both causal and disease-modifying variants, thus unlocking the potential of WGS for PID 297 diagnosis. Improved analysis methodology and better integration of parallel datasets, such as GWAS and 298 cell surface or metabolic immunophenotyping, will allow further exploration of the non-coding space 299 and enhance diagnostic yield. Such an approach promises to transform our understanding of genotype-300 phenotype relationships in PID and related immune-mediated conditions, and could redefine the clinical boundaries of immunodeficiency, add to our understanding of human immunology, and ultimately 301 302 improve patient outcomes.

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306 Acknowledgements: Funding for the NIHR-BioResource was provided by the National Institute for Health Research (NIHR, grant number RG65966). We gratefully acknowledge the participation of all NIHR 307 BioResource volunteers, and thank the NIHR BioResource centre and staff for their contribution. J.E.D.T. 308 is supported by the MRC (RG95376 and MR/L006197/1). AJT is supported by the Wellcome Trust 309 310 (104807/Z/14/Z) and the NIHR Biomedical Research Centre at Great Ormond Street Hospital for Children NHS Foundation Trust and University College London. KGCS is supported by the Medical Research 311 312 Council (program grant MR/L019027) and is a Wellcome Investigator. AJC was supported by the Wellcome [091157/Z/10/Z], [107212/Z/15/Z], [100140/Z/12/Z], [203141/Z/16/Z]; JDRF [9-2011-253], [5-313 314 SRA-2015-130-A-N]; NIHR Oxford Biomedical Research Centre and the NIHR Cambridge Biomedical Research Centre. EE has received funding from the European Union Seventh Framework Programme 315 316 (FP7-PEOPLE-2013-COFUND) under grant agreement no 609020- Scientia Fellows. 317

Author Contributions: JEDT, ES, JS, ZZ, WR, NSG, PT, AJC carried out experiments. HLA, OSB, JEDT, JHRF, DG, IS, CP, SVVD, ASJ, JM, JS, PAL, AGL, KM, EE, DE, SFJ, THK, ET performed computational analysis of the data. HLA, IS, CP, MB, CS, RL, PJRM, JS, KES conducted sample and data processing. JEDT, ES, WR, MJT, RBS, PG, HEB, AW, SH, RL, MSB, KCG, DSK, SS, SOB, TWK, WHO, AJT recruited patients, provided clinical phenotype data and confirmed genetic diagnosis. All authors contributed to the analysis of the presented results. KGCS, JEDT, HLA and OSB wrote the paper with input from all other authors. KGCS, WHO, AJT and TWK conceived and oversaw the research programme

- 324 WHO, AJT and TWK conceived and oversaw the research programme.
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327 Members of the NBR-RD PID Consortium: Zoe Adhya, Hana Alachkar, Ariharan Anantharachagan, 328 Richard Antrobus, Gururaj Arumugakani, Chiara Bacchelli, Helen E Baxendale, Claire Bethune, Shahnaz Bibi, Barbara Boardman, Claire Booth, Matthew Brown, Michael J Browning, Mary Brownlie, Matthew S 329 330 Buckland, Siobhan O Burns, Oliver S Burren, Anita Chandra, Hayley Clifford, Nichola Cooper, Godelieve J 331 de Bree, E Graham Davies, Sarah Deacock, John Dempster, Lisa A Devlin, Elizabeth Drewe, J David M 332 Edgar, William Egner, Tariq El-Shanawany, James H R Farmery, H Bobby Gaspar, Rohit Ghurye, Kimberly 333 C Gilmour, Sarah Goddard, Pavels Gordins, Sofia Grigoriadou, Scott J Hackett, Rosie Hague, Lorraine 334 Harper, Grant Hayman, Archana Herwadkar, Stephen Hughes, Aarnoud P Huissoon, Stephen Jolles, Julie 335 Jones, Yousuf M Karim, Peter Kelleher, Sorena Kiani, Nigel Klein, Taco W Kuijpers, Dinakantha S 336 Kumararatne, James Laffan, Hana Lango Allen, Sara E Lear, Hilary Longhurst, Lorena E Lorenzo, Paul A 337 Lyons, Jesmeen Maimaris, Ania Manson, Elizabeth M McDermott, Hazel Millar, Anoop Mistry, Valerie 338 Morrisson, Sai H K Murng, Iman Nasir, Sergey Nejentsev, Sadia Noorani, Eric Oksenhendler, Mark J 339 Ponsford, Waseem Qasim, Ellen Quinn, Isabella Quinti, Alex Richter, Crina Samarghitean, Ravishankar B 340 Sargur, Sinisa Savic, Suranjith L Seneviratne, W A Carrock Sewell, Fiona Shackley, Ilenia Simeoni, Kenneth 341 G C Smith, Emily Staples, Hans Stauss, Cathal L Steele, James E Thaventhiran, David C Thomas, Moira J 342 Thomas, Adrian J Thrasher, John A Todd, Anton T J Tool, Rafal D Urniaz, Steven B Welch, Lisa Willcocks, 343 Sarita Workman, Austen Worth, Nigel Yeatman, Patrick F K Yong

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- 346 The authors declare no competing financial interests.
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434

#### 435 Figure Legends

#### 436

437 Figure 1. Description of the immunodeficiency cohort and disease associations in coding regions. (a) 438 Number of index cases recruited under different phenotypic categories (red – adult cases, blue – 439 paediatric cases). (b) Number of index cases with malignancy, autoimmunity and CD4+ lymphopenia. 440 (black bar - total number of cases, blue bar - number of cases with AD-PID phenotype). (c) Number of 441 patients with reported genetic findings subdivided by gene. Previously reported variants are those 442 identified as immune disease-causing in the HGMD-Pro database. (d) Pie charts showing proportions of 443 the germline p.Arg328\* stop-gain variant and different somatic reversions in FACS-sorted blood cell populations from a male adult patient with an inherited IL2RG mutation that causes X-linked infantile 444 445 fatality. (e) BeviMed assessment of enrichment for candidate disease-causing variants in individual genes, 446 in the PID cohort relative to the rest of NBR-RD cohort. The top candidate genes (with BeviMed 447 PPA>=0.18) are shown. Named genes are those in which the variants driving the association have been 448 confirmed to be causal.

449

450 Figure 2. Assessment of WGS data for regulatory region deletions that impact upon PID (a) Schematic 451 overview of configurations of large deletions and putative damaging variants that could lead to gene loss of function. (b) Flow-chart demonstrating filtering steps to prioritise patients with candidate compound 452 453 heterozygous causal variants comprising of a rare (gnomAD v1 AF<0.001) damaging (CADD>20) coding 454 variant within a known PID gene, and a structural deletion event (with internal MAF<0.03) over the gene's 455 regulatory region. (c) Genomic configuration of the ARPC1B gene locus highlighting the compound 456 heterozygous gene variants. ExAC shows that the non-coding deletion is outside of the exome-targeted 457 regions. (d) Pedigree of patient in (c) and co-segregation of ARPC1B genotype (wt – wild-type, del – 458 deletion, fs – frameshift). (e) ARPC1A and ARPC1B protein levels in neutrophils and platelets in the patient 459 depicted in (c). (f) Histogram showing ARPC1B mRNA levels in patient depicted in (c), her sibling 460 highlighted in (d), and healthy control. (g) Allele-specific expression assay showing the ratio of wt, del and 461 fs alleles in genomic DNA (gDNA) from peripheral blood mononuclear cells of the patient and sibling. (h) 462 Relative expression of ARPC1B mRNA from each allele in the patient and sibling. Allele-specific expression 463 assessed in complementary DNA (cDNA; synthesized from pre-mRNA).

464

465 Figure 3. Telomere lengths calculated from whole-genome data can be used to identify causal rare and 466 common genomic variants associated with telomere variation. (a) Telomerecat calculated telomere 467 lengths (TLs) against age and sex in 3,313 NBR-RD recruited subjects. The Boxplot summarises the distribution of TLs within an age and gender bin; the lower, mid and upper box bounds represent the first, 468 469 second (median) and third quartile respectively. Lines extend to 1.5 times the interquartile range, and 470 outliers are marked as individual points. (b) Centiles of telomere lengths against age in PID cases. Symbols 471 represent subjects with rare genomic homozygous/hemizygous single nucleotide variants (SNV) in TERT and DKC1. (c) Top: Pedigree of individuals with DKC1 variants showing co-segregation with disease 472 473 phenotypes. The four individuals assayed by Flow-FISH are marked by dotted line. Bottom: Flow-FISH 474 assessment of telomere length in DKC1 variant carrying siblings and their spouses in granulocytes and 475 lymphocytes.

476

### 477 Figure 4. Antibody deficiency (AD-PID) GWAS identifies common variants that mediate disease risk

478 **and suggests novel monogenic candidate genes. (a)** A composite Manhattan plot for the AD-PID GWAS.

479 Blue – common variants (MAF>0.05) analysed in this study (NBR-RD) only, red – meta-analysed with

480 data from Li et al.; and purple – genome-wide significant low frequency (0.005<MAF<0.05) variants in 481 TNFRSF13B locus. Loci of interest are labelled with putative causal protein coding gene names. (b) Protein modelling of two independent MHC locus signals: residue E71 on HLA-DRB1\*1501 and residue 482 N114 on HLA-B\*0801 using PDB 1BX2 and PDB 4QRQ respectively. Protein is depicted in white, 483 484 highlighted residue in red, and peptide is in green. (c) Immune mediated trait enrichment of AD-PID association signals. CAD – coronary artery disease, CRO – Crohn's disease, RA – rheumatoid arthritis, 485 486 SLE – systemic lupus erythematosus, T1D – type 1 diabetes, T2D – type 2 diabetes and UC – ulcerative 487 colitis (See Extended Data Table 1). (d) COGS prioritisation scores of candidate monogenic causes of 488 PID using previous autoimmune targeted genotyping studies (See Supplementary Table 6) across 489 suggestive AD-PID loci (n=4). For clarity, only diseases prioritising one or more genes are shown. CEL – coeliac disease, CRO- Crohn's disease, UC – ulcerative colitis, MS – multiple sclerosis, PBC – primary 490 491 biliary cirrhosis and T1D – type 1 diabetes (e) T cells from the SOCS1 mutation patient and healthy 492 control were cultured following TCR/CD28 stimulation in the presence of anti-IFN-y and anti-IFN-yR 493 antibodies. At day 4 post-stimulation cells were washed and re-cultured without IFN-y blockade. At day 494 6 cells were stimulated for 2 hours with IFN-y and protein-lysates assessed for the indicated protein 495 expression. (Left) Representative western blot. (Right) The pSTAT1 and SOCS1 levels calculated from 496 image quantification of the western blots in 4 replicate samples. Error bars represent standard error of mean. (f) The pedigree of the CVID patient identified with a premature stop mutation in PTPN2. Carriers 497 498 of the rs2847297-G risk allele are indicated. (g) Simplified model of how SOCS1 and TC-PTP limit the 499 phosphorylated-STAT1 triggered by interferon signalling. (h) T cells from the indicated members of the 500 PTPN2 pedigree, 3 healthy controls, the SOCS1 mutation patient and a STAT1 gain of function (GOF) 501 patient were cultured for 4 days and treated +/- IFN-y for 2 hours and protein-lysates assessed for 502 protein levels. (Left) PTPN2 protein levels normalised to Tublin level (loading control). (Right) pSTAT1 503 protein levels normalised to total STAT1 level. (i) Relative expression from each allele of the PTPN2 rs2847297 locus in the sibling II.3 of the CVID patient II.1 in (f). Shown are the proportion of directly 504 genotyped individual bacterial colonies, transformed with the PCR product containing the rheumatoid 505 506 arthritis risk allele rs2847297-G generated from either gDNA or cDNA.

507

#### 508 Methods

509

#### 510 PID cohort

511 The PID patients and their family members were recruited by specialists in clinical immunology across 26 512 hospitals in the UK, and one each from the Netherlands, France and Germany. The recruitment criteria

- 513 were intentionally broad, and included the following: clinical diagnosis of common variable
- 514 immunodeficiency disorder (CVID) according to internationally established criteria (**Supplementary**
- **Table 1**); extreme autoimmunity; or recurrent and/or unusual severe infections suggestive of defective
- 516 innate or cell-mediated immunity. Patients with known secondary immunodeficiencies caused by cancer
- or HIV infection were excluded. Although screening for more common and obvious genetic causes of PID
- 518 prior to enrolment into this WGS study was encouraged, it was not a requirement. Consequently, a
- 519 minority of patients (16%) had some prior genetic testing, from single gene Sanger sequencing or MLPA520 to a gene panel screen.
- 521 To expedite recruitment a minimal clinical dataset was required for enrolment, though more detail was
- often provided. There was a large variety in patients' phenotypes, from simple "chest infections" to
- 523 complex syndromic features, and the collected phenotypic data of the sequenced individuals ranged
- 524 from assigned disease category only to detailed clinical synopsis and immunophenotyping data. The
- 525 clinical subsets used to subdivide PID patients were based on ESID definitions, as shown in
- 526 Supplementary Table 1.
- 527 To facilitate analysis by grouping patients with a degree of phenotypic coherence while excluding some 528 distinct and very rare clinical subtypes of PID that may have different aetiologies, a group of patients
- 529 was determined to have antibody deficiency-associated PID (AD-PID). This group comprised 733 of the
- 530 886 unrelated index cases, and included all patients with CID, CVID or Antibody Defect ticked on the
- recruitment form, together with patients requiring IgG replacement therapy and those with specified
- 532 low levels of IgG/A/M. SCID patients satisfying these criteria were not assigned to the AD-PID cohort.
- 533

### 534 WGS data processing

535 Details of DNA sample processing, whole genome sequencing, data processing pipeline, quality checks, 536 alignment and variant calling, ancestry and relatedness estimation, variant normalisation and 537 annotation, large deletion calling and filtering, and allele frequency calculations, are fully described in 538 [NIHR BioResource, in preparation; see Cover Letter]. Briefly, DNA or whole blood EDTA samples were 539 processed and quality checked according to standard laboratory practices and shipped on dry ice to the 540 sequencing provider (Illumina Inc, Great Chesterford, UK). Illumina Inc performed further QC array 541 genotyping, before fragmenting the samples to 450bp fragments and processing with the Illumina 542 TruSeq DNA PCR-Free Sample Preparation kit (Illumina Inc., San Diego, CA, USA). Over the three-year 543 duration of the sequencing phase of the project, different instruments and read lengths were used: for 544 each sample, either 100bp reads on three HiSeq2500 lanes; or 125bp reads on two HiSeq2500 lanes; or 545 150bp reads on a single HiSeq X lane. Each delivered genome had a minimum 15X coverage over at least 546 95% of the reference autosomes. Illumina performed the alignment to GRCh37 genome build and 547 SNV/InDel calling using their Isaac software, while large deletions were called with their Manta and 548 Canvas algorithms. The WGS data files were received at the University of Cambridge High Performance

- 549 Computing Service (HPC) for further QC and processing by our Pipeline team.
- 550 For each sample, we estimated the sex karyotype and computed pair-wise kinship coefficients using
- 551 PLINK, which allowed us to identify sample swaps and unintended duplicates, assign ethnicities,
- 552 generate networks of closely related individuals (sometimes undeclared relatives from across different
- 553 disease domains) and a maximal unrelated sample set (for the purposes of allele frequency estimation
- and control dataset in case-control analyses). Variants in the gVCF files were normalised and loaded into

- 555 an HBase database, where Overall Pass Rate (OPR) was computed within each of the three read length 556 batches, and the lowest of these OPR values (minOPR) assigned to each variant.
- 557 Large deletions were merged and analysed collectively, as described in [NIHR BioResource, in
- 558 preparation]. The analyses presented here are based on SNVs/InDels with OPR>0.98, and a set of
- deletions found through the SVH method to have high specificity after extensive manual inspection of
- 560 individual deletion calls. Variants were annotated with Sequence Ontology terms according to their
- 561 predicted consequences, their frequencies in other genomic databases (gnomAD, UK10K, 1000
- 562 Genomes), if they have been associated with a disease according to the HGMD Pro database, and
- 563 internal metrics (AN, AC, AF, OPR).
- 564

### 565 Diagnostic reporting

566 We screened all genes in the IUIS 2015 classification for potentially causal variants. SNVs and small 567 InDels were filtered based on the following criteria: OPR>0.95; having a protein-truncating consequence,

- 568 gnomAD AF<0.001 and internal AF<0.01; or present in the HGMD Pro database as DM variant. Large
- 569 deletions called by both Canvas and Manta algorithms, passing standard Illumina quality filters,
- 570 overlapping at least one exon, and classified as rare by the SVH method were included in the analysis. In
- order to aid variant interpretation and consistency in reporting, phenotypes were translated into Human
- 572 Phenotype Ontology (HPO) terms as much as possible. Multi-Disciplinary Team (MDT) then reviewed
- each variant for evidence of pathogenicity and contribution to the phenotype, and classified them
- according to the American College of Medical Genetics (ACMG) guidelines<sup>38</sup>. Only variants classified as
- 575 Pathogenic or Likely Pathogenic were systematically reported, but individual rare (gnomAD AF<0.001) or
- 576 novel missense variants that BeviMed analysis (see below) highlighted as having a posterior probability 577 of pathogenicity >0.2 were additionally considered as Variants of Unknown Significance (VUS). If the
- 578 MDT decided that they were likely to be pathogenic and contribute to the phenotype, they were also
- 579 reported and counted towards the overall diagnostic yield. All variants and breakpoints of large
- 580 deletions reported in this study were confirmed by Sanger sequencing using standard protocols.
- 581

### 582 <u>BeviMed</u>

583 We used BeviMed<sup>1</sup> to evaluate the evidence for association between case/control status and rare

- variant allele counts in each gene. We inferred a posterior probability of association (PPA) under
- 585 Mendelian inheritance models (dominant and recessive), and different variant selection criteria
- 586 ("moderate" and "high" impact variants based on functional consequences predicted by the Variant
- 587 Effect Predictor<sup>39</sup>). All genes were assigned the same prior probability of association with the disease of
- 588 0.01, regardless of their previously published associations with an immune deficiency phenotype. Genes
- 589 for which BeviMed inferred a PPA to be >=0.18 when summed over all four combinations of inheritance
- 590 model and variant selection criteria (each configuration being given a prior probability of association of
- 591 0.0025) are shown in **Fig. 1f**. Given each of the association models, the posterior probability that each 592 variant is pathogenic is also computed. We used a variant-level posterior probability of pathogenicity
- 593 >0.2 to select potentially pathogenic missense variants in known PID genes to report back.
- 594

# 595 <u>Telomerecat</u>

596 Average telomere length was calculated from whole-genome sequence data using Telomerecat, as 597 reported previously<sup>22</sup>. Batch differences caused by changes in sequencing platform differences were

598 normalised by using a linear model. The linear model was defined as:

$$length = \beta_0 + \beta_1 batch_2 + \beta_2 batch_3 \dots \beta_N batch_N + \epsilon$$

599

- 600 where  $\beta$  are regression coefficients, and batch represents a dummy variable denoting the plate a sample
- was sequenced on. For each plate the relevant coefficient was subtracted from all of the observedtelomere lengths within each plate.
- After adjusting for batch effects, telomere length was compared to age in 3,313 NBR-RD subjects. We
- obtained a strong negative correlation with age (r = -0.56, Pearson's correlation), thus validating
- Telomerecat as a reliable method for estimating telomere lengths. We found that each year of
- additional age was equivalent to a 33bp deterioration in telomere length (Supplementary Fig. 6).
- 607 Although this observed negative correlation is well established within the literature, we obtain a
- particularly high correlation with our method, which could be partly driven by the wide age range of oursample set.
- To normalise telomere lengths for comparison of samples from disparate age and gender, the followinglinear model was fitted to the data using age as a continuous variable and gender as a dummy variable:

$$length = \beta_0 + \beta_1 age + \beta_2 age^2 + \beta_3 age^3 + \beta_4 gender + e$$

612

The relevant residuals produced by the cubic model were subtracted from the mean telomere length ofthe cohort. These adjusted telomere lengths were used in the GWAS analysis.

- To assess for monogenic causes of telomere shortening, subjects were identified within the PID cohort
- that had telomere lengths below the 10<sup>th</sup> centile of age adjusted values and had hemizygous or
- 617 homozygous SNVs that occurred gnomAD AF<0.001 in *TERC, TERT, NHP2, TINF2, NOP10, PARN, ACD,*
- 618 WRAP53, CTC1, RTEL1 or DKC1 genes.
- 619

### 620 <u>AD-PID GWAS</u>

- 621 GWAS was performed both on the whole PID cohort (N cases = 886) and on a subset of AD-PID cases (N
- 622 cases = 733); here we present the results of the latter analysis, which was cleaner and less noisy despite 623 a reduced sample size. We used 9225 unrelated samples from non-PID NBR-RD cohorts as controls.
- 624 Variants were selected from a merged VCF file were filtered to include bi-allelic SNPs with overall
- 625 MAF>=0.05 and minOPR=1 (100% pass rate). We ran PLINK logistic association test under an additive
- 626 model using the read length, sex, and first 10 principal components from the ethnicity analysis as
- 627 covariates. After filtering out SNPs with HWE  $p < 10^{-6}$ , we were left with the total of 4,993,945 analysed
- 628 SNPs. There was minimal genomic inflation of the test statistic (lambda = 1.027), suggesting population
- 629 substructure and sample relatedness had been appropriately accounted for. The only genome-wide
- 630 significant (p<5x10<sup>-8</sup>) signal was at the MHC locus, with several suggestive (p<1x10<sup>-5</sup>) signals
- 631 (Supplementary Fig. 7). We repeated the analysis with more relaxed SNP filtering criteria using
- 632 MAF>=0.005 and minOPR>0.95. The only additional signal identified were the three *TNFRSF13B* variants
- 633 shown in **Extended Data Fig. 1**.
- 634 We obtained summary statistics data from the Li et al. CVID Immunochip case-control study<sup>9</sup> and
- 635 performed a fixed effects meta-analysis on 95,417 variants shared with our AD-PID GWAS. For each of
- the genome-wide and suggestive loci after meta-analysis, we conditioned on the lead SNP by including it
- as an additional covariate in the logistic regression model, to determine if the signal is driven by the
- 638 single or multiple hits at those loci. Only the MHC locus showed evidence of multiple independent
- 639 signals (Supplementary Fig. 8).
- 640

### 641 MHC locus imputation

- 642 We imputed classical HLA alleles using the method implemented in the SNP2HLA v1.0.3 package<sup>40</sup>,
- 643 which uses Beagle v3.0.4 for imputation and the HapMap CEU reference panel. We imputed allele

- dosages and best-guess genotypes of 2-digit and 4-digit classical HLA alleles, as well as amino acids of
- the MHC locus genes *HLA-A*, *HLA-B*, *HLA-C*, *HLA-DRB1*, *HLA-DQA1* and *HLA-DQB1*. We tested the
- association of both allele dosages and genotypes using the logistic regression implemented in PLINK,
- and obtained similar results. We then used the best-guess genotypes to perform the conditional analysis
- 648 in PLINK, since conditioning is not implemented in a model with allele dosages.
- 649

## 650 Allele Specific Expression

651 RNA and gDNA were extracted from PBMCs using the AllPrep kit (Qiagen) as per the manufacturer's

- 652 instructions. RNA was reverse transcribed to make cDNA using the SuperScript<sup>™</sup> VILO<sup>™</sup> cDNA synthesis
- 653 kit with appropriate minus reverse transcriptase controls, as per the manufacturer's instructions. The
- region of interest in the gDNA and 1:10 diluted cDNA was amplified using Phusion (Thermo Fisher) and
- the following primers on a G-Storm thermal cycler with 30 seconds at 98°C then 35 cycles of 98°C 10
- 656 seconds, 60°C 30 seconds, 72°C 15 seconds.

# 657 **ARPC1B**

658 The region of interest spanning the frameshift variant was amplified using the following primers:

- 659 Forward: GGGTACATGGCGTCTGTTTC / Reverse: CACCAGGCTGTTGTCTGTGA
- 660 PCR products were run on a 3.5% agarose gel. Bands were cut out and product extracted using the QIA
- 661 Quick Gel Extraction Kit (Qiagen), as per protocol. Expected products were confirmed by Sanger
- sequencing. 4ul fresh PCR product was used in a TOPO<sup>®</sup> cloning reaction (Invitrogen) and used to
- transform One Shot<sup>™</sup> TOP10 chemically competent E. coli. These were cultured overnight then spread
- 664 on LB agar plates. Individual colonies were picked and genotyped. ARPC1B mRNA expression was
- assessed using a Taqman gene expression assay with 18S and EEF1A1 as control genes. Each sample was
- run in triplicate for each gene with a no template control. PCR was run on a LightCycler<sup>®</sup> (Roche) with 2
- 667 mins 50°C, 20 seconds  $95^{\circ}$ C then 45 cycles of  $95^{\circ}$ C 3 seconds,  $60^{\circ}$ C 30 seconds.
- 668 *PTPN2*
- 669 PTPN2 ASE protocol is modified from above. RNA and genomic DNA were extracted from PBMCs using
- 670 the AllPrep Kit (Qiagen). RNA was treated with Turbo DNAse (Thermo) and reverse transcribed to
- 671 generate cDNA using the SuperScript IV VILO master mix (Thermo). The intronic region of interest in
- 672 gDNA and cDNA was amplified by two nested PCR reactions using Phusion enzyme (Thermo). The
- 673 primers (F1/R1) and nested primers (F2/R2) used were:
- 674 Forward\_1: aaagtctggagcaggcaggg / Reverse\_1: tgggggaactggttatgctttc
- 675 Forward\_2: ggagctatgatcacgccacatg / Reverse\_2: atgctttctggttgggctgac
- 676
- 677 PCR products were run on a 1% agarose gel. Bands were cut out and product extracted using the QIA
- 678 Quick Gel Extraction Kit (Qiagen), as per protocol. Expected products were confirmed by Sanger
- sequencing. 5ng fresh PCR product was used in a TOPO<sup>®</sup> cloning reaction (Invitrogen) and used to
- transform One Shot<sup>™</sup> TOP10 chemically competent E. coli. These were cultured overnight then spread
- on LB agar plates. Individual colonies were picked and genotyped. PTPN2 mRNA expression was
- assessed using a Taqman SNP genotyping assay and on a LightCycler (Roche).
- 683
- 684 PAGE and Western Blot analysis
- 685 Samples were separated by SDS polyacrylamide gel electrophoresis and transferred onto a nitrocellulose
- 686 membrane. Individual proteins were detected with antibodies against ARPC1b (goat polyclonal
- antibodies, ThermoScientific, Rockford, IL, USA), against ARPC1a (rabbit polyclonal antibodies, Sigma, St
- 688 Louis, USA) and against actin (mouse monoclonal antibody, Sigma). Secondary antibodies were either
- 689 donkey-anti-goat-IgG IRDye 800CW, Goat-anti-mouse-IgG IRDye 800CW or Donkey-anti-rabbit-IgG IRDye

690 680CW (LI-COR Biosciences, Lincoln, NE, USA). Quantification of bound antibodies was performed on an 691 Odyssey Infrared Imaging system (LI-COR Biosciences, Lincoln, NE, USA).

692

#### 693 Phasing of SOCS1 variants

694 To phase common rs2286974 variant with the novel stop-gain SOCS1 variant (chr16:11348854 695 T>TGCGGC) identified in the same patient, we performed long-read WGS with Oxford Nanopore 696 Technologies (ONT). The sample was prepared using the 1D ligation library prep kit (SQK-LSK108), and 697 genomic libraries were sequenced on R9.4 flowcells. Sequencing was carried out on GridION system, 698 read sequences were extracted from base-called FAST5 files by Guppy (v0.5.1) to generate FASTQ files, which were then aligned against the GRCh37/hg19 human reference genome using minimap2 (v2.2). 699 700 Four runs were performed in order to reach an average coverage of 14x, with a median read length of 701 5006 ± 3981. Haplotyping and genotyping was performed with MarginPhase.

702

### 703 Structural deletion analysis

704 Structural (length >50bp) deletions (MAF>0.03) were called as previously described<sup>41</sup>. For all

downstream analysis we used gencode v26 annotations downloaded from

706 [ftp://ftp.sanger.ac.uk/pub/gencode/Gencode\_human/release\_26/GRCh37\_mapping/gencode.v26lift37

notation.gtf.gz]. We defined promoters as a window +/- 500bp of any protein coding gene

transcriptional start site (TSS). In order to associate cis regulatory elements (cRE) with putative target

genes we combined by physical location overlap, super enhancer cRE annotations from <sup>18</sup>, with

710 promoter capture Hi-C (pcHi-C) from <sup>19</sup>, matching by tissue. We next computed the overlap of structural 711 variants occurring in the PID cohort with cREs for which putative target genes were available. We

712 classified overlaps between deletions and functional annotations into three non-mutually exclusive

categories; `prom' - overlaps focal gene promoter, `exon' - overlaps focal gene exon, `pse' - overlaps

714 Hnisz *et al.*<sup>18</sup> SE annotation linked to focal gene by pcHi-C. We compiled a catalogue of compound

heterozygous deletions where there was evidence in the same individual for a damaging (CADD>20) rare

- 716 (gnomAD AF<0.001) variant within the same gene.
- 717

# 718 AD-PID GWAS Enrichment

719 Due to the size of the AD-PID cohort, we were unable to use LD-score regression<sup>42</sup> to assess genetic

correlation between distinct and related traits. We therefore adapted the previous enrichment method

<sup>721</sup> `blockshifter`<sup>43</sup> in order to assess evidence for the enrichment of AD-PID association signals in a

compendium of 9 GWAS European Ancestry summary statistics was assembled from publicly available

data. We removed the MHC region from all downstream analysis [GRCh37 chr6:25-45Mb]. To adjust for

724 linkage disequilibrium (LD), we split the genome into 1cM recombination blocks based on HapMap

recombination frequencies <sup>44</sup>. For a given GWAS trait, for n variants within LD block b we used

726 Wakefield's synthesis of asymptotic Bayes factors (aBF)<sup>45</sup> to compute the posterior probability that the

727  $i^{th}$  variant is causal (*PPCV<sub>i</sub>*) under single causal variant assumptions<sup>46</sup> :

728

729

$$PPCV_i = \frac{aBF_i\pi_i}{\sum_{j=1}^n (aBF_j\pi_j) + 1}$$

Here  $\pi_i = \pi_j$  are flat prior probabilities for a randomly selected variant from the genome to be causal

and we use the value  $1 \times 10^{-447}$ . We sum over these PPCV within an LD block, b to obtain the posterior

732 probability that *b* contains a single causal variant (PPCB).

- To compute enrichment for trait *t*, we convert PPCBs into a binary label by applying a threshold such
- that  $PPCB_t > 0.95$ . We apply these block labels for trait t, to PPCBs (computed as described above) for
- our AD-PID cohort GWAS, using them to compute a non-parametric Wilcoxon rank sum statistic, W
- representing the enrichment. Whilst the aBF approach naturally adjusts for LD within a block, residual
- LD between blocks may exist. In order to adjust for this and other confounders (e.g. block size) we use a
- circularised permutation technique<sup>48</sup> to compute W<sub>null</sub>. To do this, for a given chromosome, we select
   recombination blocks, and circularise such that beginning of the first block adjoins the end of the last.
- Permutation proceeds by rotating the block labels, but maintaining AD-PID PPCB assignment. In this way
- 741 many permutations of W<sub>null</sub> can be computed whilst conserving the overall block structure.
- For each trait we used 10<sup>4</sup> permutations to compute adjusted Wilcoxon rank sum scores using *wgsea*
- 743 [https://github.com/chr1swallace/wgsea] R package.
- 744

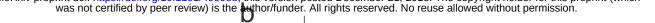
#### 745 PID monogenic candidate gene prioritisation

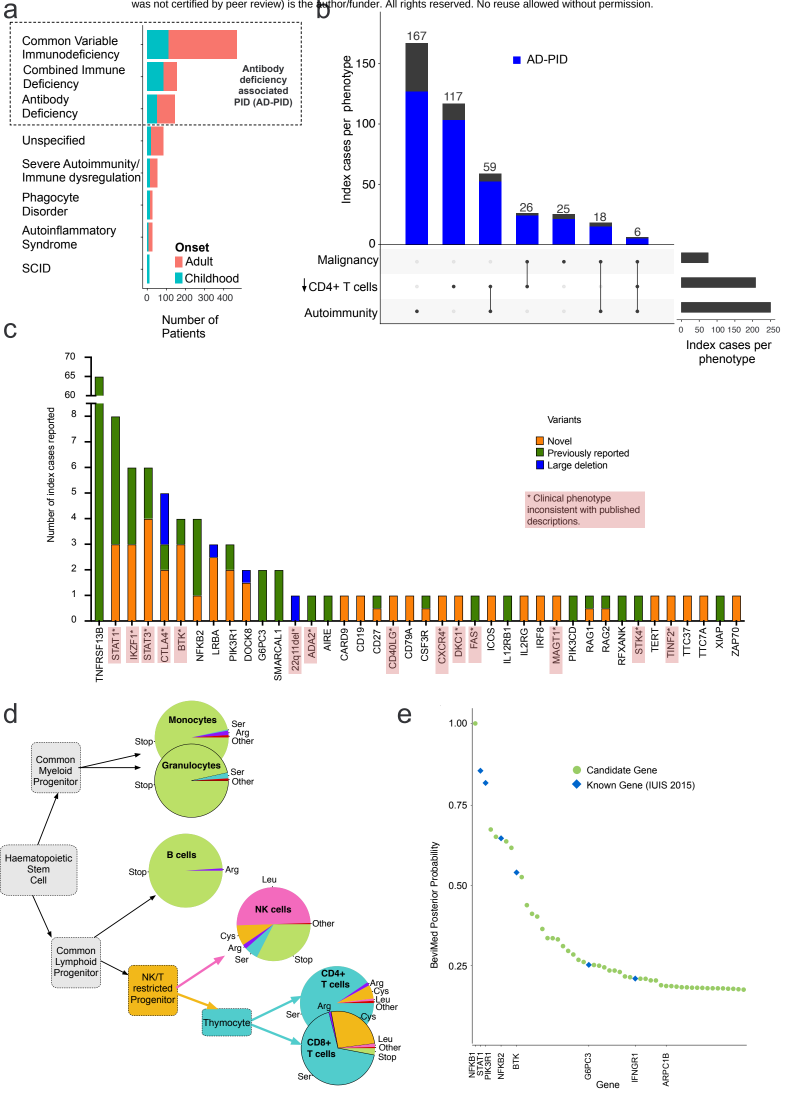
- 746 We hypothesised, given the genetic overlap with antibody associated PID, that common regulatory
- variation, elucidated through association studies of immune-mediated disease, might prioritise genes
- harbouring damaging LOF variants underlying PID. Firstly, using summary statistics from our combined
- fixed effect meta-analysis of AD-PID, we compiled a list of densely genotyped ImmunoChip regions
- containing one or more variant where  $P<1x10^{-5}$ . Next, we downloaded ImmunoChip (IC) summary
- statistics from ImmunoBase (accessed 30/07/2018) for all 11 available studies. For each study we
- intersected PID suggestive regions, and used COGS (https://github.com/ollyburren/rCOGS) in
- conjunction with promoter-capture Hi-C datasets for 17 primary cell lines<sup>19,43</sup> in order to prioritise genes.
- 754 We filtered by COGS score to select genes with a COGS score >0.5<sup>19,43</sup> to obtain a list of 11 protein
- 755 coding genes.
- 756 We further hypothesised that genes harbouring rare LOF variation causal for PID would be intolerant to
- variation. We thus downloaded pLI scores<sup>49</sup> and took the product between these and the COGS scores
- to compute an `overall' prioritisation score across each trait and gene combination. We applied a final
- filter taking forward only those genes having an above average `overall' score to obtain a final list of 6
   candidate genes (Fig. 4d). Finally, we filtered the cohort for damaging rare (gnomAD AF<0.001) protein-</li>
- 761 truncating variants (frameshift, splice-site, nonsense) within these genes in order to identify individuals
- 762 for functional follow up.
- 763
- 764 <u>Statistical analysis</u>
- 765 Statistical analysis was carried out using R (3.3.3 "Another Canoe") and Graphpad Prism (version 7)
- 766 unless otherwise stated. R code for running major analyses are available at
- 767 https://github.com/ollyburren/pid\_thaventhiran\_et\_al.
- 768

# 769 Methods References

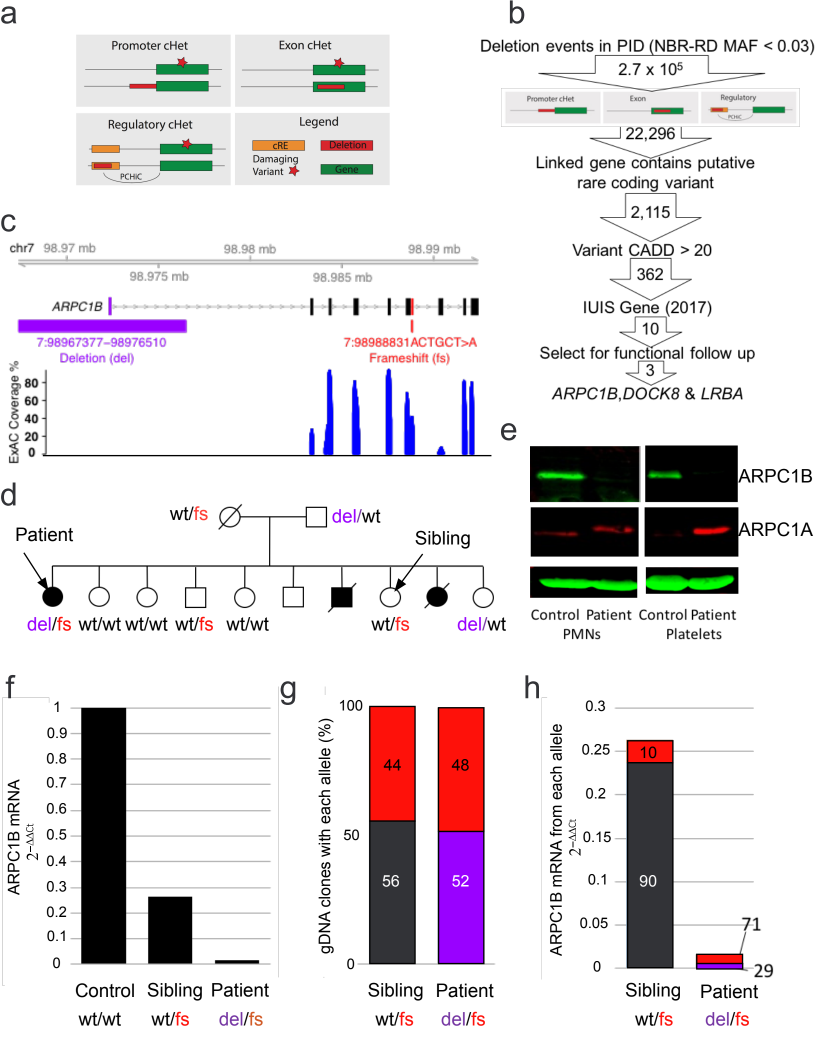
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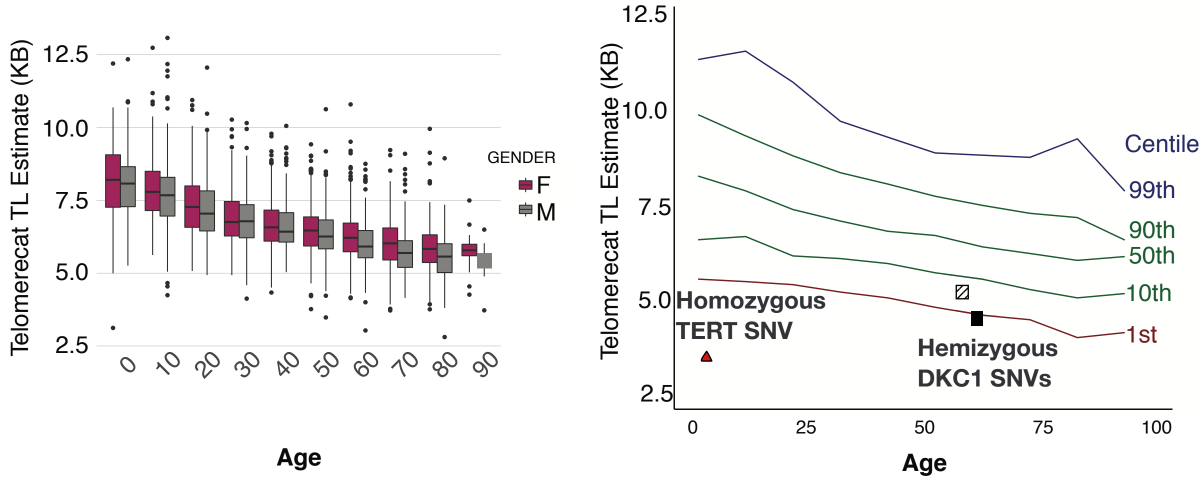




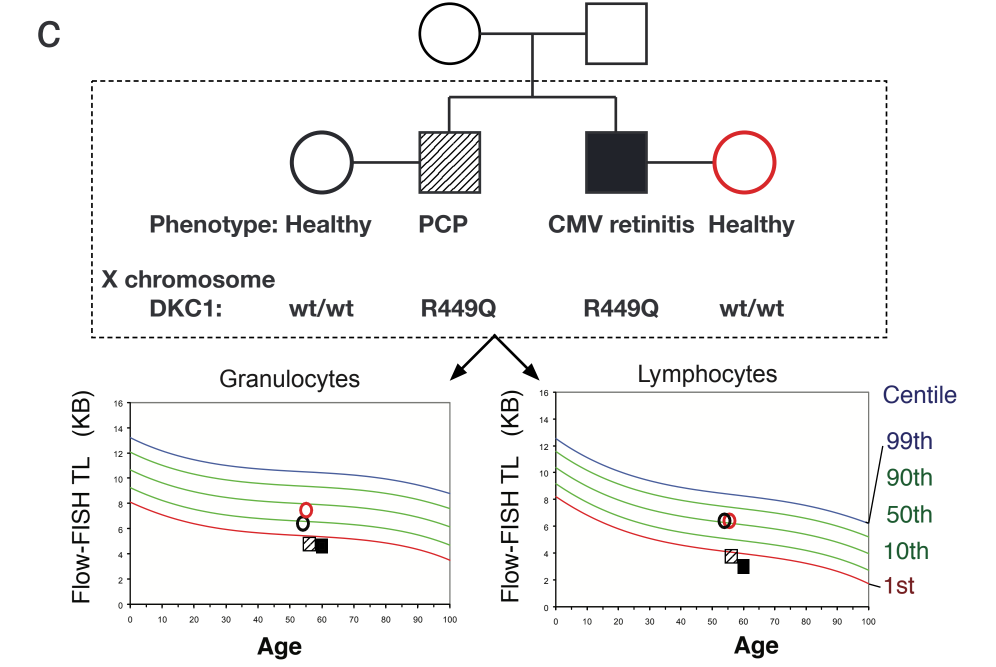
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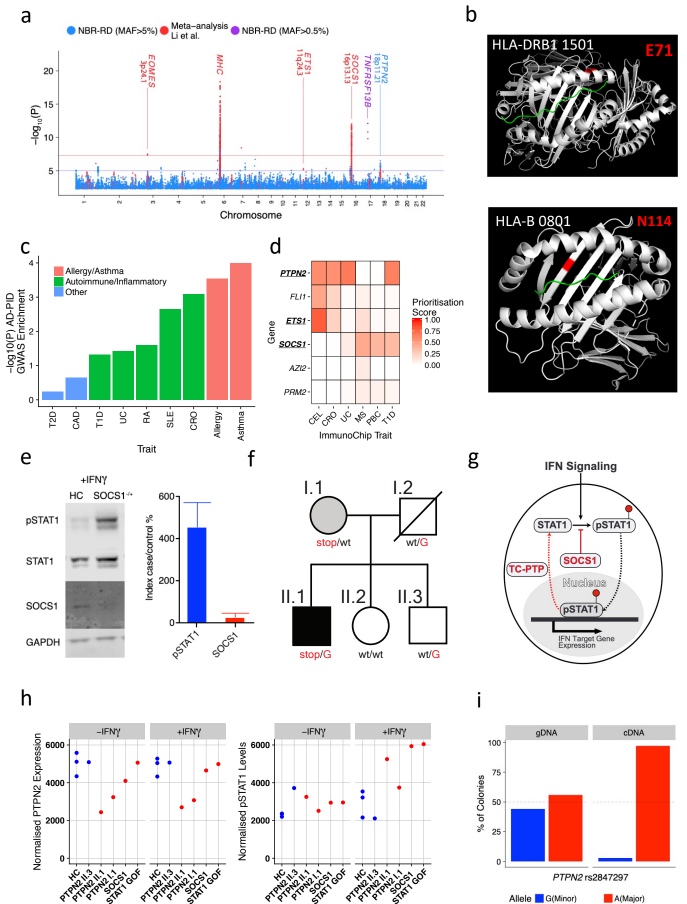


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D





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Allele G(Minor)

A(Major)

# **Extended Data Items Legends**

Extended Data Table 1. GWAS studies used in enrichment analysis

**Extended Data Fig. 1.** Regional AD-PID association plot of 17p11.2 (*TNFRSF13B/TACI*) region. Tracks are as follows: **AD-PID -log10(P)** dot plot of AD-PID association, index SNP ( is purple, others are coloured based on LD information from UK10K project with red indicating high LD (r^2>0.9), blue low (r^2<0.2) and grey where no information available. **Gene** - Cannonical gene annotation (Ensembl V75), **GWAS** - location of index variants from other immune-mediated disease, **CD4**, **B**, **Mon** putative regulatory regions in CD+T-cells, Total B cells and Monocytes computed from the union of ATAC-Seq and H3K27ac ChIP-Seq data, **pcHi-C** - Promoter Capture Hi-C interactions, in above primary cell types.

**Extended Data Fig. 2.** Regional AD-PID meta-analysis association plot of 3p24.1 (*EOMES*) region. Tracks as described for Extended Data Fig.1 with the exception of **Meta -log(P)** which shows dot plot of AD-PID association meta-analysis with Li *et al.* Detail shows location of RA index SNP that overlaps AD-PID index variant and its promoter proximity.

**Extended Data Fig. 3.** Comparison of GWAS association signals at 18p11.21 for Li *et al.*, **NBR-RD AD-PID** (this study), and **Meta** (Meta-analysis). Y-axis is -log10(P) of univariate association statistic. Top SNP in each study is marked in cyan.

**Extended Data Fig. 4.** Locus plot of Regional AD-PID meta-analysis association plot of 16p13.13 (*CLEC16A/SOCS1*). Tracks as described in Extended Data Fig. 2.

