

雑誌

発表者氏名	論文タイトル名	発表誌	巻号	ページ	出版年
Suzuki, E., et al. (共著者11名中8番目)	De novo frameshift mutation in fibroblast growth factor 8 in a male patient with gonadotropin deficiency	<i>Horm. Res. Paediatr.</i>	81	139-144	2014
Kubo, A., et al. (共著者27名中21番目)	Mutations in SERPINB7, encoding a member of the serine protease inhibitor superfamily, cause Nagashima-type palmoplantar keratosis	<i>Am. J. Hum. Genet.</i>	93	945-956	2013
Fukami, M., et al. (共著者15名中10番目)	Genomic basis of aromatase excess syndrome: recombination-and replication-mediated rearrangements leading to CYP19A1 overexpression	<i>J. Clin. Endocrinol. Metab.</i>	98	2013-2021	2013

共著者が10名を超える場合は、筆頭著者名 *et al.* で記載

書籍

著者氏名	論文タイトル名	書籍全体の編集者名	書籍名	出版社名	出版地	出版年	ページ
Miyake, N., Kosho, T., Matsumoto, N.	Ehlers-Danlos syndrome associated with glycosaminoglycan abnormalities.	J. Halper	<i>Progress in Heritable Soft Connective Tissue Diseases, Advances in Experimental Medicine and Biology</i> 802	Springer, Netherland	Houten, Netherland	2014	145-159

雑誌

発表者氏名	論文タイトル名	発表誌	巻号	ページ	出版年
Gupta, V.A. et al. (共著者29名中26番目)	Identification of KLHL41 mutations implicates BTB-Kelch-mediated ubiquitination as an alternate pathway to myofibrillar disruption in nemaline myopathy	<i>Am. J. Hum. Genet.</i>	93	1108-1017	2013
Nishiguchi, K.M. et al. (共著者22名中18番目)	Whole genome sequencing in patients with retinitis pigmentosa reveals pathogenic DNA structural changes and NEK2 as a new disease gene	<i>Proc. Natl. Acad. Sci. USA</i>	110	16139-16144	2013
Nakamura K, et al. (共著者26名中25番目)	De novo mutations in GNAO1 encoding a Gao subunit of heterotrimeric G proteins, cause epileptic encephalopathy	<i>Am. J. Hum. Genet.</i>	93	496-505	2013
Ravenscroft, G. et al (共著者55名中54番目)	Mutations in KLHL40 are a frequent cause of severe autosomal-recessive nemaline myopathy	<i>Am. J. Hum. Genet.</i>	93	6-18	2013
Nakajima, M. et al. (共著者31名中28番目)	Mutations in B3GALT6, which encodes a glycosaminoglycan linker region enzyme, cause a spectrum of skeletal and connective tissue disorders	<i>Am. J. Hum. Genet.</i>	92	927-934	2013
Saito, H. et al. (共著者21名中21番目)	De novo mutations in the autophagy gene WDR45 cause static encephalopathy of childhood with neurodegeneration in adulthood	<i>Nat. Genet.</i>	45	445-449	2013

共著者が10名を超える場合は、筆頭著者名 et al. で記載

雑誌

発表者氏名	論文タイトル名	発表誌	巻号	ページ	出版年
Hamaguchi, Y. et al. (共著者15名中12番目)	Common and distinct clinical features in adult patients with anti-aminoacyl-tRNA synthetase antibodies: heterogeneity within the syndrome	<i>PLoS One</i>	8	e60442	2013
Terao, C. et al. (共著者18名中14番目)	Three groups in the 28 Joints for rheumatoid arthritis synovitis - analysis using more than 17,000 assessments in the KURAMA database	<i>PLoS One</i>	8	e59341	2013
Terao, C. et al. (共著者32名中28番目)	Two susceptibility loci to Takayasu arteritis reveal a synergistic role of the IL12B and HLA-B regions in a Japanese population	<i>Am. J. Hum. Genet.</i>	93	289-297	2013

共著者が10名を超える場合は、筆頭著者名 *et al.* で記載

研究分担者：日笠 幸一郎

書籍

著者氏名	論文タイトル名	書籍全体の編集者名	書籍名	出版社名	出版地	出版年	ページ
日笠 幸一郎、 松田 文彦	日本人の遺伝 子リファレン スライブライ ーデータベー ス	松本 直通	エクソーム解析 —成果と将来	医歯薬出版 株式会社	東京	2013	352-357

雑誌

発表者氏名	論文タイトル名	発表誌	巻号	ページ	出版年
Yamamoto, H. et al. (共著者15名中2番目)	Novel germline mutation in the transmembrane domain of HER2 in familial lung adenocarcinomas	<i>J. Natl. Cancer Inst.</i>	106	djt338	2014
Takahashi, Y. et al. (共著者40名中9番目)	ERBB4 mutations that disrupt the neuregulin-ErbB4 pathway cause amyotrophic lateral sclerosis type 19	<i>Am. J. Hum. Genet.</i>	93	900-905	2013

共著者が 10 名を超える場合は、筆頭著者名 *et al.* で記載

研究分担者：寺尾 知可史

雑誌

発表者氏名	論文タイトル名	発表誌	巻号	ページ	出版年
Okada, Y. et al. (共著者96/94名中5番目)	Genetics of rheumatoid arthritis contributes to biology and drug discovery	Nature	506	376-381	2014
Terao, C. et al. (共著者32名中1番目)	Two susceptibility loci to Takayasu arteritis reveal a synergistic role of the IL12B and HLA-B regions in a Japanese population	Am. J. Hum. Genet.	93	289-297	2013
Terao, C. et al. (共著者17名中1番目)	Association of Takayasu arteritis with HLA-B*67:01 and two amino acids in HLA-B protein	Rheumatol.	52	1769-1774	2013

共著者が 10 名を超える場合は、筆頭著者名 *et al.* で記載

VII. 学会発表に関する一覧表

研究代表者：松田 文彦

発表者名	演題名	学会名	会場	日時
Matsuda, F.	Genomic analysis of immune-related diseases using GWAS and WES	The Second Kyoto Symposium on Bioinformatics for Next Generation Sequencing with Applications in Human Genetics	Kyoto, Japan	March 14, 2014
Matsuda, F.	Genome-wide association studies in IgG4-RD	Second International Symposium on IgG4 and Related Diseases	Honolulu, USA	February 18, 2014
Matsuda, F.	The Nagahama Study as a model for the comprehensive human bioscience	Kyoto Symposium on Bioinformatics for Next Generation Sequencing with Application in Human Genetics	Kyoto, Japan	January 19, 2013
松田 文彦	地域に根ざした新時代の予防医学の試み	2013 年度ソニー／医科歯科クリニックサミット	東京医科大学 鈴木記念講堂（東京）	2013 年 8 月 20 日
松田 文彦	ヒト生命情報統合研究とゲノムコホート	第 20 回日本遺伝子診療学会大会教育講演	アクトシティ浜松コングレスセンター（浜松）	2013 年 7 月 20 日
松田 文彦	難病・がん等の疾患分野の医療の実用化研究事業の成果を基にした原因遺伝子変異データベースの構築	日本予防医学協会 医療と健康のシンポジウム	全国社会保険協会連合会研修センター（東京）	2013 年 3 月 20 日
松田 文彦	ヒト生命情報統合研究とその情報基盤	大阪大学蛋白研究所セミナー「ビッグデータ時代に向けた医療データベース」	大阪大学中之島センター（大阪）	2013 年 3 月 8 日
松田 文彦	分子を通して自分の健康をながめる～ながはま 0 次コホート研究と次世代の予防医療～	いわて東北メディカル・メガバンク機構 発足記念シンポジウム	岩手医科大学 矢巾キャンパス（矢巾）	2013 年 2 月 2 日

研究分担者：辻 省次

発表者名	演題名	学会名	会場	日時
三井 純、辻 省次	ペーソナルゲノム解析が医療を変貌させる	第 85 回日本遺伝学会市民公開講座	慶應義塾大学日吉キャンパス(横浜)	2013 年 9 月 21 日
三井 純、後藤 順、辻 省次	エクソーム解析による遺伝的異質性の高い疾患に対する遺伝子検査	第 20 回日本遺伝子診療学会	アクシティ浜松(浜松)	2013 年 7 月 19 日

研究分担者：松原 洋一

発表者名	演題名	学会名	会場	日時
Izumi, R. et al.	A mutation in A-band titin is associated with hereditary myopathy with early respiratory failure in a Japanese family	American Society of Human Genetics 63rd Annual Meeting	Boston, USA	October 22-26, 2013
Niihori, T. et al.	Exome sequencing identifies mutations in a novel gene in patients with Noonan syndrome	American Society of Human Genetics 63rd Annual Meeting	Boston, USA	October 22-26, 2013
Aoki, Y., Niihori, T., Inoue, S. and Matsubara, Y.	Genetic syndromes associated with the Ras/MAPK pathway and the identification of mutations in a new gene, RIT1, for Noonan syndrome	Third International Meeting on Genetic Syndromes of the Ras/MAPK Pathway: Towards a Therapeutic Approach	Orland, USA	August 2-4, 2013
新堀 哲也ら	エクソームシークエンスによる Noonan 症候群新規原因遺伝子 RIT1 の同定	日本人類遺伝学会 第 58 回大会	江陽グランドホテル(仙台)	2013 年 11 月 20~23 日
青木 洋子ら	次世代シークエンサーを用いたヌーナン症候群の遺伝子診断と新規原因遺伝子検索	日本人類遺伝学会 第 58 回大会	江陽グランドホテル(仙台)	2013 年 11 月 20~23 日
緒方 勤ら	エクソーム解析により TBX1 変異が同定された家族性の特徴的顔貌・鼻咽頭閉鎖不全・低 Ca 血症を呈する 5 例	日本人類遺伝学会 第 58 回大会	江陽グランドホテル(仙台)	2013 年 11 月 20~23 日
井泉 瑠美子ら	Myofibrillar myopathy の大家系における次世代型シークエンサーを用いた原因遺伝子の同定	第 54 回日本神経学会学術大会	東京国際フォーラム(東京)	2013 年 5 月 29 日

研究分担者：松本 直通

発表者名	演題名	学会名	会場	日時
Matsumoto, N., et al.	De novo mutations in the autophagy gene encoding WDR45 (WIPI4) cause static encephalopathy of childhood with neurodegeneration in adulthood (oral)	European Conference of Human Genetic 2013	Paris, France	June 9, 2013
Matsumoto, N.	Mendelian exome analysis (oral)	The 10 th International Workshop on Advanced Genomics	Tokyo, Japan	May 21, 2013
松本 直通	次世代シーケンサー解析と小児医療	平成 25 年度第 2 回バイオビジネス・スタートアップへゲノム解析による疾患の原因究明とその基盤技術～	AP 横浜西口（横浜）	2014 年 3 月 5 日
松本 直通	次世代シーケンサーによる難病遺伝子解析（教育講演）	第 8 回ファブリー病シンポジウム	東京コンファレンスセンター有明（東京）	2014 年 3 月 1 日
松本 直通	WDR45 変異が来すヒト疾患	大阪大学蛋白質研究所セミナー・オートファジーと疾患	大阪大学蛋白質研究所（吹田）	2014 年 2 月 21 日
松本 直通	希少疾患・難病の全エクソーム解析 —現状と課題—	日経バイオテク「希少疾患・難病の治療薬開発におけるゲノム活用」	秋葉原コンベンションホール（東京）	2013 年 12 月 3 日
松本 直通	ヒト疾患エクソーム解析の現状と課題（シンポジスト）	第 58 回日本人類遺伝学会大会	江陽グランドホテル（仙台）	2013 年 11 月 23 日
松本 直通	次世代シーケンサーを用いた疾患ゲノム解析：現状と限界（特別講演）	第 22 回発達腎研究会	高槻市生涯学習センター（高槻）	2013 年 9 月 27 日
松本 直通	NGS がもたらしたヒト疾患ゲノム解析のパラダイムシフト（基調講演）	現場の会第三回研究会	神戸国際会議場（神戸）	2013 年 9 月 4 日
松本 直通	疾患ゲノム解析における次世代シーケンサーの有用性（シンポジスト）	第 20 回日本遺伝子診療学会大会	アクシティ－浜松コングレスセンター（浜松）	2013 年 7 月 19 日
松本 直通	次世代シーケンサーを用いてわかつてきしたこと（特別講演）	第 17 回小児分子内分泌研究会	札幌北広島クラッセホテル（札幌）	2013 年 7 月 7 日

研究分担者：山田 亮

発表者名	演題名	学会名	会場	日時
Narahara, M., Matsuda, F., Yamada, R., et al.	Establishing an eQTL map of the Japanese population.	American Society of Human Genetics 63rd Annual Meeting	Boston, USA	October 25, 2013
Narahara, M., Matsuda, F., Yamada, R., et al	日本人における eQTL マップの構築	日本人類遺伝学会 第 58 回大会	江陽グランドホテル(仙台)	2013 年 11 月 22 日

研究分担者：日笠 幸一郎

発表者名	演題名	学会名	会場	日時
日笠 幸一郎	日本人の遺伝子リファレンスライブラリーデータベース	日本人類遺伝学会 第 58 回大会	江陽グランドホテル(仙台)	2013 年 11 月 21 日
日笠 幸一郎	大規模ゲノムコホート研究に基づく日本人遺伝子多型データベース	医学研究のためのバイオデータベース講習会	東海大学医学部伊勢原キャンパス講堂 B(伊勢原)	2013 年 12 月 25 日
日笠 幸一郎	ヒト生命情報統合研究に向けた大規模ゲノムコホート事業の推進	NGS 現場の会	神戸国際会議場(神戸)	2013 年 9 月 4 日

研究分担者：寺尾 知可史

発表者名	演題名	学会名	会場	日時
Yoshifuji, H., Terao, C., Murakami, K., Kawabata, D., Ohmura, K., Fujii, T., Kawaguchi, Y., Yamanaka, H., Mimori, T.	Association between HLA-B's amino acid variation and disease-susceptibility to Takayasu arteritis.	American College of Rheumatology 2013	San Diego, USA	October 27, 2013
寺尾 知可史、吉藤元、木村 彰方、松村貴由、大村 浩一郎、成瀬 妙子、佐藤 愛子、前島 康浩、和田庸子、成田 一衛、川口 鎮司、山中 寿、前川 平、小川 誠司、小室 一成、永井 良三、田原 康玄、磯部光章、三森 経世、松田 文彦	IL12B は高安動脈炎の新規疾患感受性遺伝子であり HLA-B*52:01 と相互作用を示す	第 41 回日本臨床免疫学会	海峡メッセ(下関)	2013 年 11 月 28 日
寺尾 知可史、吉藤元、村上 孝作、川端大介、大村 浩一郎、秋月 正史、川口 鎮司、山中 寿、三森 経世	高安動脈炎感受性と相關する 3 つの新規 HLA-B アミノ酸多型	第 57 回日本リウマチ学会総会・学術集会	国立京都国際会館(京都)	2013 年 4 月 20 日

VIII. 研究成果による特許等の知的財産権の 出願・登録状況

研究分担者：松本 直通

1. 特願 2013-123660 才津浩智／松本直通 小児期のてんかんおよび不随意運動をきたす疾患の検出方法 2013年6月12日
2. 特願 2013-157339 松本直通／三宅紀子 ケトン血症を伴うリーコンドリア複合体III欠乏症患者または保因者の検出法 2013年7月31日
3. 特願 2013-252720 鶴崎美徳／松本直通 Coffin-Siris 症候群の新規遺伝子診断法 2013年12月6日
4. PCT/JP2013/71620 松本直通／三宅紀子 ミトコンドリア複合体III欠乏症患者又は保因者の検出方法 2013年8月9日・2014年2月7日

IX. 研究成果の刊行物・別刷

論文リスト

1. Okada, Y., Wu, D., Trynka, G., Raj, T., Terao, C., Ikari, K., Kochi, Y., Ohmura, K., Suzuki, A., Yoshida, S., Graham, R. R., Manoharan, A., Ortmann, W., Bhagale, T., Denny, J. C., Carroll, R. J., Eyler, A. E., Greenberg, J. D., Kremer, J. M., Pappas, D. A., Jiang, L., Yin, J., Ye, L., Su, D. F., Yang, J., Xie, G., Keystone, E., Westra, H. J., Esko, T., Metspalu, A., Zhou, X., Gupta, N., Mirel, D., Stahl, E. A., Diogo, D., Cui, J., Liao, K., Guo, M. H., Myouzen, K., Kawaguchi, T., Coenen, M. J. H., van Riel, P. L. C. M., van de Laar, M. A. F. J., Guchelaar, H. J., Huizinga, T. W. J., Dieude, P., Mariette, X., Bridges Jr, S. L., Zhernakova, A., Toes, R. E. M., Tak, P. P., Miceli-Richard, C., Bang, S. Y., Lee, H. S., Martin, J., Gonzalez-Gay, M. A., Rodriguez-RodriguezL., Rantapaa-Dahlqvist, S., Arlestig, L., Choi, H. K., Kamatani, Y., Galan, P., Lathrop, M., the RACI consortium, the GARNET consortium, Eyre, S., Bowes, J., Barton, A., de Vries, N., Moreland, L. W., Criswell, L. A., Karlson, E. W., Taniguchi, A., Yamada, R., Kubo, M., Liu, J. S., Bae, S. C., Worthington, J., Padyukov, L., Klareskog, L., Gregersen, P. K., Raychaudhuri, S., Stranger, B. E., De Jager, P. L., Franke, L., Visscher, P. M., Brown, M. A., Yamanaka, H., Mimori, T., Takahashi, A., Xu, H., Behrens, T. W., Siminovitch, K. A., Momohara, S., Matsuda, F., Yamamoto, K. and Plenge, R. M. (2014) Genetics of rheumatoid arthritis contributes to biology and drug discovery. *Nature* 506, 376-381.
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3. Yamamoto, H., Higasa, K., Sakaguchi, M., Shien, K., Soh, J., Ichimura, K., Furukawa, M., Hashida, S., Tsukuda, K., Takigawa, N., Matsuo, K., Kiura, K., Miyoshi, S., Matsuda, F. and Toyooka, S. (2013) Novel germline mutation in the transmembrane domain of *HER2* in familial lung adenocarcinomas. *J. Natl. Cancer Inst.* Dec 7. [Epub ahead of print]
4. Oishi, M., Yamashiro, K., Miyake, M., Akagi-Kurashige, Y., Kumagai, K., Nakata, I., Nakanishi, H., Yoshikawa, M., Oishi, A., Gotoh, N., Tsujikawa, A., Yamada, R., Matsuda, F. and Yoshimura, N. (2013) Association between *ZIC2*, *RASGRF1*, and *SHISA6* genes and high myopia in Japanese subjects. *Invest. Ophthalmol. Vis. Sci.* 54, 7492-7497.
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Genetics of rheumatoid arthritis contributes to biology and drug discovery

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A major challenge in human genetics is to devise a systematic strategy to integrate disease-associated variants with diverse genomic and biological data sets to provide insight into disease pathogenesis and guide drug discovery for complex traits such as rheumatoid arthritis (RA)¹. Here we performed a genome-wide association study meta-analysis in a total of >100,000 subjects of European and Asian ancestries (29,880 RA cases and 73,758 controls), by evaluating ~10 million single-nucleotide polymorphisms. We discovered 42 novel RA risk loci at a genome-wide level of significance, bringing the total to 101 (refs 2–4). We devised an *in silico* pipeline using established bioinformatics methods based on functional annotation⁵, *cis*-acting expression quantitative trait loci⁶ and pathway analyses^{7–9}—as well as novel methods based on genetic overlap with human primary immunodeficiency, haematological cancer somatic mutations and knockout mouse phenotypes—to identify 98 biological candidate genes at these 101 risk loci. We demonstrate that these genes are the targets of approved therapies for RA, and further suggest that drugs approved for other indications may be repurposed for the treatment of RA. Together, this comprehensive genetic study sheds light on fundamental genes, pathways and cell types that contribute to RA pathogenesis, and provides empirical evidence that the genetics of RA can provide important information for drug discovery.

We conducted a three-stage trans-ethnic meta-analysis (Extended Data Fig. 1). On the basis of the polygenic architecture of RA¹⁰ and shared genetic risk among different ancestry^{3,4}, we proposed that combining a genome-wide association study (GWAS) of European and Asian ancestry would increase power to detect novel risk loci. In stage 1, we combined 22 GWAS for 19,234 cases and 61,565 controls of European and Asian ancestry^{2–4}. We performed trans-ethnic, European-specific and Asian-specific GWAS meta-analysis by evaluating ~10 million single-nucleotide polymorphisms (SNPs)¹¹. Characteristics of the cohorts, genotyping platforms and quality control criteria are described in Extended Data Table 1 (overall genomic control inflation factor $\lambda_{GC} < 1.075$).

Stage 1 meta-analysis identified 57 loci that satisfied a genome-wide significance threshold of $P < 5.0 \times 10^{-8}$, including 17 novel loci (Extended Data Fig. 2). We then conducted a two-step replication study (stage 2 for *in silico* and stage 3 for *de novo*) in 10,646 RA cases and 12,193 controls for the loci with $P < 5.0 \times 10^{-6}$ in stage 1. In a combined analysis of stages 1–3, we identified 42 novel loci with $P < 5.0 \times 10^{-8}$ in any of the trans-ethnic, European or Asian meta-analyses. This increases the total number of RA risk loci to 101 (Table 1 and Supplementary Table 1).

Comparison of 101 RA risk loci revealed significant correlations of risk allele frequencies (RAFs) and odds ratios (ORs) between Europeans and Asians (Extended Data Fig. 3a–c; Spearman's $\rho = 0.67$ for RAF and 0.76 for OR; $P < 1.0 \times 10^{-13}$), although five loci demonstrated population-specific associations ($P < 5.0 \times 10^{-8}$ in one population but $P > 0.05$ in the other population without overlap of the 95% confidence intervals (95% CIs) of the ORs). In the population-specific genetic risk model, the 100 RA risk loci outside of the major histocompatibility complex (MHC) region¹² explained 5.5% and 4.7% of heritability in Europeans and Asians, respectively, with 1.6% of the heritability explained by the novel loci. The trans-ethnic genetic risk model, based on the RAF from

one population but the OR from the other population, could explain the majority (>80%) of the known heritability in each population (4.7% for Europeans and 3.8% for Asians). These observations support our hypothesis that the genetic risk of RA is shared, in general, among Asians and Europeans.

We assessed enrichment of 100 non-MHC RA risk loci in epigenetic chromatin marks¹³ (Extended Data Fig. 3d). Of 34 cell types investigated, we observed significant enrichment of RA risk alleles with trimethylation of histone H3 at lysine 4 (H3K4me3) peaks in primary CD4⁺ regulatory T cells (T_{reg} cells; $P < 1.0 \times 10^{-5}$). For the RA risk loci enriched with T_{reg} H3K4me3 peaks, we incorporated the epigenetic annotations along with trans-ethnic differences in patterns of linkage disequilibrium to fine-map putative causal risk alleles (Extended Data Fig. 3e, f).

We found that approximately two-thirds of RA risk loci demonstrated pleiotropy with other human phenotypes (Extended Data Fig. 4), including immune-related diseases (for example, vitiligo, primary biliary cirrhosis), inflammation-related or haematological biomarkers (for example, fibrinogen, neutrophil counts) and other complex traits (for example, cardiovascular diseases).

Each of 100 non-MHC RA risk loci contains on average ~4 genes in the region of linkage disequilibrium (in total 377 genes). To prioritize systematically the most likely biological candidate gene, we devised an *in silico* bioinformatics pipeline. In addition to the published methods that integrate data across associated loci^{7,8}, we evaluated several biological data sets to test for enrichment of RA risk genes, which helps to pinpoint a specific gene in each loci (Extended Data Figs 5, 6 and Supplementary Tables 2–4).

We first conducted functional annotation of RA risk SNPs. Sixteen per cent of SNPs were in linkage disequilibrium with missense SNPs ($r^2 > 0.80$; Extended Data Fig. 5a, b). The proportion of missense RA risk SNPs was higher compared with a set of genome-wide common SNPs (8.0%), and relatively much higher in the explained heritability (~26.8%). Using *cis*-acting expression quantitative trait loci (*cis*-eQTL) data obtained from peripheral blood mononuclear cells (5,311 individuals)⁶ and from CD4⁺ T cells and CD14⁺CD16[−] monocytes (212 individuals), we found that RA risk SNPs in 44 loci showed *cis*-eQTL effects (false discovery rate (FDR) q or permutation $P < 0.05$; Extended Data Table 2).

Second, we evaluated whether genes from RA risk loci overlapped with human primary immunodeficiency (PID) genes¹⁴, and observed significant overlap (14/194 = 7.2%, $P = 1.2 \times 10^{-4}$; Fig. 1a and Extended Data Fig. 5c). Classification categories of PID genes showed different patterns of overlap; the highest proportion of overlap was in ‘immune dysregulation’ (4/21 = 19.0%, $P = 0.0033$) but there was no overlap in ‘innate immunity’.

Third, we evaluated overlap with cancer somatic mutation genes¹⁵, under the hypothesis that genes with cell growth advantages may contribute to RA development. Among 444 genes with registered cancer somatic mutations¹⁵, we observed significant overlap with genes implicated in haematological cancers (17/251 = 6.8%, $P = 1.2 \times 10^{-4}$; Fig. 1b and Extended Data Fig. 5d), but not with genes implicated in non-haematological cancers (6/221 = 2.7%, $P = 0.56$).

Table 1 | Novel rheumatoid arthritis risk loci identified by trans-ethnic GWAS meta-analysis in >100,000 subjects

SNP	Chr	Genes	A1/A2 (+)	Trans-ethnic			European			Asian	
				OR (95% CI)	P	OR (95% CI)	P	OR (95% CI)	P	OR (95% CI)	P
rs227163	1	TNFRSF9	C/T	1.04 (1.02–1.06)	3.9×10^{-4}	1.00 (0.97–1.03)	9.3×10^{-1}	1.11 (1.08–1.16)*	3.1×10^{-9} *		
rs28411352	1	MTF1-INPP5B	T/C	1.11 (1.08–1.14)*	2.8×10^{-12} *	1.10 (1.07–1.14)*	5.9×10^{-9} *	1.12 (1.06–1.19)	7.8×10^{-5}		
rs2105325	1	LOC100506023	C/A	1.12 (1.08–1.15)*	6.9×10^{-13} *	1.12 (1.08–1.15)*	3.3×10^{-11} *	1.13 (1.04–1.23)	5.2×10^{-3}		
rs10175798	2	LBH	A/G	1.08 (1.06–1.11)*	1.1×10^{-9} *	1.09 (1.06–1.12)*	4.2×10^{-8} *	1.07 (1.02–1.13)	6.4×10^{-3}		
rs6732565	2	ACOXL	A/G	1.07 (1.05–1.10)*	2.7×10^{-8} *	1.10 (1.07–1.14)*	9.4×10^{-9} *	1.04 (1.00–1.08)	4.0×10^{-2}		
rs6715284	2	CFLAR-CASP8	G/C	1.15 (1.10–1.20)*	1.8×10^{-9} *	1.15 (1.10–1.20)*	2.5×10^{-9} *				
rs4452313	3	PLCL2	T/A	1.09 (1.06–1.12)*	1.6×10^{-10}	1.11 (1.08–1.15)*	5.2×10^{-11} *	1.04 (0.99–1.09)	9.2×10^{-2}		
rs3806624	3	EOMES	G/A	1.08 (1.05–1.11)*	8.6×10^{-9} *	1.08 (1.05–1.12)*	2.8×10^{-8} *	1.06 (0.99–1.14)	1.0×10^{-1}		
rs9826828	3	IL20RB	A/G	1.44 (1.28–1.61)*	8.6×10^{-10}	1.44 (1.28–1.61)*	8.7×10^{-10} *				
rs13142500	4	CLNK	C/T	1.10 (1.07–1.13)*	3.0×10^{-9} *	1.10 (1.06–1.15)	2.4×10^{-6}	1.10 (1.04–1.15)	2.8×10^{-4}		
rs2664035	4	TEC	A/G	1.07 (1.04–1.10)	9.5×10^{-8}	1.08 (1.05–1.11)*	3.3×10^{-8} *	1.03 (0.97–1.08)	3.3×10^{-1}		
rs9378815	6	IRF4	C/G	1.09 (1.06–1.12)*	1.7×10^{-10}	1.09 (1.05–1.12)	1.4×10^{-7}	1.10 (1.04–1.15)	2.3×10^{-4}		
rs2234067	6	ETV7	C/A	1.15 (1.10–1.20)*	1.6×10^{-9} *	1.14 (1.09–1.19)*	4.1×10^{-8} *	1.22 (1.06–1.41)	7.0×10^{-3}		
rs9373594	6	PPI4	T/C	1.09 (1.06–1.12)*	3.0×10^{-9} *	1.07 (1.02–1.12)	6.5×10^{-3}	1.11 (1.07–1.15)*	4.8×10^{-8} *		
rs67250450	7	JAZF1	T/C	1.10 (1.07–1.14)*	3.7×10^{-9} *	1.11 (1.07–1.14)*	2.6×10^{-9} *	1.02 (0.84–1.23)	8.5×10^{-1}		
rs4272	7	CDK6	G/A	1.10 (1.06–1.13)*	5.0×10^{-9} *	1.10 (1.07–1.14)*	1.2×10^{-8} *	1.06 (0.98–1.15)	1.3×10^{-1}		
rs998731	8	TPD52	T/C	1.08 (1.05–1.11)*	1.9×10^{-8} *	1.09 (1.06–1.12)*	6.6×10^{-9} *	1.02 (0.96–1.10)	4.9×10^{-1}		
rs678347	8	GRHL2	G/A	1.08 (1.05–1.11)*	1.6×10^{-8} *	1.10 (1.06–1.13)*	7.3×10^{-9} *	1.03 (0.98–1.10)	2.6×10^{-1}		
rs1516971	8	PVT1	T/C	1.15 (1.10–1.20)*	1.3×10^{-10}	1.16 (1.11–1.21)*	3.2×10^{-11} *				
rs12413578	10	10p14	C/T	1.20 (1.13–1.29)*	4.8×10^{-8} *	1.20 (1.12–1.29)	7.5×10^{-8}				
rs793108	10	ZNF438	T/C	1.08 (1.05–1.10)*	1.3×10^{-9} *	1.07 (1.04–1.10)	6.1×10^{-7}	1.09 (1.04–1.14)	4.4×10^{-4}		
rs2671692	10	WDFY4	A/G	1.07 (1.05–1.10)*	2.8×10^{-9} *	1.06 (1.03–1.09)	2.6×10^{-5}	1.10 (1.05–1.14)	9.9×10^{-5}		
rs726288	10	SFTP4	T/C	1.14 (1.07–1.20)	1.6×10^{-5}	0.96 (0.86–1.06)	4.1×10^{-1}	1.22 (1.14–1.31)*	8.8×10^{-9} *		
rs968567	11	FADS1-FADS2-FADS3	C/T	1.12 (1.07–1.16)*	1.8×10^{-8} *	1.12 (1.07–1.16)*	1.8×10^{-8} *				
rs4409785	11	CEP57	C/T	1.12 (1.09–1.16)*	1.2×10^{-11} *	1.12 (1.08–1.16)*	3.6×10^{-9} *	1.16 (1.07–1.27)	4.3×10^{-4}		
chr11:107967350	11	ATM	A/G	1.21 (1.13–1.29)*	1.4×10^{-8} *	1.21 (1.13–1.29)*	1.1×10^{-8} *				
rs73013527	11	ETS1	C/T	1.09 (1.06–1.12)*	1.2×10^{-10} *	1.08 (1.05–1.11)	1.0×10^{-6}	1.14 (1.08–1.21)	4.1×10^{-6}		
rs773125	12	CDK2	A/G	1.09 (1.06–1.12)*	1.1×10^{-10} *	1.09 (1.06–1.12)*	2.1×10^{-8} *	1.10 (1.04–1.17)	1.1×10^{-3}		
rs10774624	12	SH2B3-PTPN11	G/A	1.09 (1.06–1.13)*	6.8×10^{-9} *	1.09 (1.06–1.13)*	6.9×10^{-9} *				
rs9603616	13	COG6	C/T	1.10 (1.07–1.13)*	1.6×10^{-12} *	1.11 (1.07–1.14)*	2.8×10^{-11} *	1.08 (1.02–1.14)	1.0×10^{-2}		
rs3783782	14	PRKCH	A/G	1.14 (1.09–1.18)*	2.2×10^{-9} *	1.12 (0.96–1.31)	1.4×10^{-1}	1.14 (1.09–1.19)*	4.4×10^{-9} *		
rs1950897	14	RAD51B	T/C	1.10 (1.07–1.13)*	8.2×10^{-11} *	1.09 (1.06–1.12)*	5.0×10^{-8} *	1.16 (1.08–1.25)	1.1×10^{-4}		
rs4780401	16	TXND11	T/G	1.07 (1.05–1.10)*	4.1×10^{-8} *	1.09 (1.06–1.13)*	8.7×10^{-9} *	1.03 (0.98–1.08)	2.5×10^{-1}		
rs72634030	17	C1QBP	A/C	1.12 (1.08–1.17)*	1.5×10^{-9} *	1.12 (1.06–1.19)	2.9×10^{-5}	1.12 (1.07–1.18)	9.6×10^{-6}		
rs1877030	17	MED1	C/T	1.09 (1.06–1.12)*	1.9×10^{-8} *	1.09 (1.05–1.13)	1.3×10^{-5}	1.09 (1.04–1.14)	3.2×10^{-4}		
rs2469434	18	CD226	C/T	1.07 (1.05–1.10)*	8.9×10^{-10} *	1.05 (1.02–1.08)	6.7×10^{-4}	1.11 (1.07–1.15)*	1.2×10^{-8} *		
chr19:10771941	19	ILF3	C/T	1.47 (1.30–1.67)*	8.6×10^{-10} *	1.47 (1.30–1.67)*	8.8×10^{-10} *				
rs73194058	21	IFNGR2	C/A	1.08 (1.05–1.12)	1.2×10^{-6}	1.13 (1.08–1.18)*	2.6×10^{-8} *	1.03 (0.98–1.08)	2.9×10^{-1}		
rs1893592	21	UBASH3A	A/C	1.11 (1.08–1.14)*	7.2×10^{-12} *	1.11 (1.07–1.15)*	9.8×10^{-9} *	1.11 (1.05–1.18)	1.3×10^{-4}		
rs11089637	22	UBE2L3-YDJC	C/T	1.08 (1.05–1.11)*	2.1×10^{-9} *	1.10 (1.06–1.15)	2.0×10^{-7}	1.06 (1.02–1.10)	8.9×10^{-4}		
rs909685	22	SYNGR1	A/T	1.13 (1.10–1.16)*	1.4×10^{-16} *	1.11 (1.08–1.15)*	6.4×10^{-12} *	1.23 (1.14–1.33)	2.0×10^{-7}		
chrX:78464616	X	P2RY10	A/C	1.11 (1.07–1.15)*	3.5×10^{-8} *	1.16 (0.78–1.75)	4.6×10^{-1}	1.11 (1.07–1.15)*	3.6×10^{-8} *		

SNPs newly associated with $P < 5.0 \times 10^{-8}$ in the combined study of the stage 1 GWAS meta-analysis and the stages 2 and 3 replication studies of trans-ethnic (Europeans and Asians), European or Asian ancestry are indicated. SNPs, positions and alleles are based on the positive (+) strand of NCBI build 37. A1 represents an RA risk allele. Chr, chromosome; OR, odds ratio; 95% CI, 95% confidence interval. Full results of the studies are available in Supplementary Table 1. Hyphens between gene names indicate that several candidate RA risk genes were included in the region.

*Association results with $P < 5.0 \times 10^{-8}$.

Fourth, we evaluated overlap with genes implicated in knockout mouse phenotypes¹⁶. Among the 30 categories of phenotypes¹⁶, we observed 3 categories significantly enriched with RA risk genes ($P < 0.05/30 = 0.0017$): ‘haematopoietic system phenotype’, ‘immune system phenotype’, and ‘cellular phenotype’ (Extended Data Fig. 5e).

Last, we conducted molecular pathway enrichment analysis (Fig. 1c and Extended Data Fig. 5f). We observed enrichment (FDR $q < 0.05$) for T-cell-related pathways, consistent with cell-specific epigenetic marks, as well as enrichment for B-cell and cytokine signalling pathways (for example, interleukin (IL)-10, interferon, granulocyte-macrophage colony-stimulating factor (GM-CSF)). For comparison, our previous RA GWAS meta-analysis² did not identify the B-cell and cytokine signalling pathways, thereby indicating that as more loci are discovered, further biological pathways are identified.

On the basis of these new findings, we adopted the following 8 criteria to prioritize each of the 377 genes from the 100 non-MHC RA risk loci (Fig. 2 and Extended Data Fig. 6a–c): (1) genes with RA risk missense variant ($n = 19$); (2) *cis*-eQTL genes ($n = 51$); (3) genes prioritized by PubMed text mining⁷ ($n = 90$); (4) genes prioritized by protein–protein interaction (PPI)⁸ ($n = 63$); (5) PID genes ($n = 15$); (6) haematological cancer somatic mutation genes ($n = 17$); (7) genes prioritized by associated knockout mouse phenotypes ($n = 86$); and (8) genes prioritized by molecular pathway analysis⁹ ($n = 35$).

Ninety-eight genes (26.0%) had a score ≥ 2 , which we defined as ‘candidate biological RA risk genes’. Nineteen loci included multiple biological RA risk genes (for example, *IL3* and *CSF2* at chromosome 5q31), whereas no biological gene was selected from 40 loci (Supplementary Table 5).

To provide empirical evidence of the pipeline, we evaluated relationships of the gene scores to independent genomic or epigenetic information. Genes with higher biological scores were more likely to be the nearest gene to the risk SNP (18.6% for gene score < 2 and 49.0% for gene score ≥ 2 ; $P = 2.1 \times 10^{-8}$), and also to be included in the region where RA risk SNPs were overlapping with H3K4me3 T_{reg} peaks (41.9% for gene score < 2 and 57.1% for gene score ≥ 2 ; $P = 0.034$). Further, T_{reg} cells demonstrated the largest increase in overlapping proportions with H3K4me3 peaks for increase of biological gene scores compared with other cell types (Extended Data Fig. 6d).

Finally, we evaluated the potential role of RA genetics in drug discovery. We proposed that if human genetics is useful for drug target validation, then it should identify existing approved drugs for RA. To test this ‘therapeutic hypothesis’¹, we obtained 871 drug target genes corresponding to approved, in clinical trials or experimental drugs for human diseases^{17,18} (Supplementary Table 6). We evaluated whether any of the protein products from the identified biological RA risk genes, or any genes from a direct PPI network with such protein products