

Supplementary Table 4. Coding variants at loci associated with RA or its subsets and their functional prediction^a

Lead variant (chr:pos)	Coding variant	R ²	EA	OA	Gene	Coding effect	Available in dbNSFP v4.1			MPC			Mutation		Mutation		BayesDel			BayesDel		fathmmxMKL	fathmmxF						
							Damaging	Tolerated	Neutral	(rankscore)	SIFT	SIFT4G	LRT	Taster	Assessor	FATHMM	PROVEAN	MetaSVM	MetaLR	MxCAP	PrimateAI	DEOGEN2	addAF	noAF	LIST-S2	Aloft	coding	coding	
chr1:2800059	chr1:295307	0.89	G	A	MMEL1	missense	x	1	9	5	0.22	T	T	N	P	N	D	N	T	T	.	T	T	T	T	.	N	N	
chr1:2800059	chr1:2596694	0.96	C	T	MMEL1	splice region	-																						
chr1:2800059	chr1:2609830	0.92	G	A	MMEL1	splice region	-																						
chr1:2800059	chr1:2800059	1.00	C	T	TTC34	missense	x	0	2	2	.	.	.	P		
chr1:113834946	chr1:113834946	1.00	G	A	PTPN22	missense	x	0	10	4	0.15	T	T	N	P	N	T	N	T	T	.	T	T	T	T	.	N	.	
chr1:161506414*	chr1:161506414	1.00	T	C	FCGR2A	stop gained	x	0	3	3	.	.	.	N	P	N	N
chr1:161506414*	chr1:161506414	1.00	G	A	FCGR2A	missense	x	0	10	4	0.26	T	T	N	P	M	T	N	T	T	.	T	T	T	T	.	N	N	
chr2:191073180**	chr2:191073180**	1.00	A	T	STAT4	missense	x	10	5	1	0.67	T	T	D	D	L	D	N	D	D	D	T	T	T	D	D	.	D	D
chr3:58197909**	chr3:58197909	1.00	A	G	DNASE1L3	missense	x	10	4	0	.	D	D	D	P	H	D	D	T	T	.	D	D	D	D	D	.	D	D
chr6:137678425#	chr6:137874929	<0.2	T	G	TNFAIP3	missense	x	2	10	2	0.47	T	T	N	P	L	T	D	T	T	.	T	T	T	T	.	D	N	
chr7:128938247	chr7:128938247	1.00	G	T	IRF5	splice donor	-																						
chr12:111446804	chr12:111446804	1.00	C	T	SHZB3	missense	x	0	10	5	0.12	T	T	N	P	N	T	N	T	T	.	T	T	T	T	.	N	N	
chr13:39771329**	chr13:39771329**	0.95	CTT	TTT	COG6	splice region	-																						
chr13:397788092*	chr13:397788092	0.95	CTT	TTT	COG6	splice region	-																						
chr14:92651884	chr14:92651884	1.00	T	C	RIN3	missense	x	4	8	2	0.37	D	D	N	P	M	T	D	T	T	.	T	T	T	T	.	D	N	
chr17:39908216*	chr17:39908216	1.00	A	G	GSDMB	missense	x	1	9	3	0.23	T	T	N	P	.	T	D	T	T	.	T	T	T	T	.	N	N	
chr17:39908216*	chr17:39908216	0.80	T	C	GSDMB	missense	x	2	8	3	0.57	T	D	N	P	.	T	D	T	T	.	T	T	T	T	.	N	N	
chr17:39908216*	chr17:39908216	1.00	C	T	GSDMB	splice acceptor	x	0	2	2	.	.	.	P	T	T	.	N	N	.	N	N	
chr19:10352442	chr19:10352442	1.00	C	G	TYK2	missense	x	7	7	1	0.66	D	D	N	D	L	T	D	T	T	.	T	T	T	D	.	D	D	
chr19:10359299**	chr19:10359299**	1.00	C	A	TYK2	missense	x	7	6	1	0.77	D	D	N	P	M	T	D	T	T	.	T	T	T	D	D	.	D	D
chr19:10354167**	chr19:10354167**	1.00	A	G	TYK2	missense	x	28	4	0	0.72	D	D	D	D	L	D	D	D	D	D	T	T	D	D	.	D	D	
#	chr19:10471767	<0.2	C	A	PDE4A	missense	x	1	9	4	.	D	T	N	P	N	T	N	T	T	.	T	T	T	T	.	N	N	
chr21:44236991	chr21:44236991	1.00	C	T	ICOSLG	missense	x	0	10	4	0.71	T	T	N	P	M	T	N	T	T	.	T	T	T	T	.	N	N	

Lead variants that are only GWAS significant in seropositive RA (*) or in RA overall (**), other variants are significant in both. Coding variants that are independent secondary signals after adjustment for lead variants are marked with (**), see Supplementary Tables 2 and 8.

^aAssociation with seropositive RA of missense variant in TNFAIP3 (chr6:137874929, OR=1.25, P=7.7x10⁻¹) is not GWAS significant after adjustment for lead signal at the locus (chr6:137678425, adjusted OR=1.18, adjusted P=3.2x10⁻³), despite the fact that they are not correlated (R²<0.2). Similarly, missense variant in PDE4A (chr19:10467167, OR=0.90, P=6.3x10⁻¹⁰)Functional prediction was performed using the dbNSFP (version 4.1) database (<https://sites.google.com/site/jpoggen/dbNSFP>).

dbNSFP results are presented as a summarized score, rankscore or as D, damaging, T, tolerated, N, neutral, indicating predicted functional effect of the sequence variant.

Sources of prediction tools used in the dbNSFP v4.1 database:

MVP 1.1, https://github.com/schenabu/misense_MVC release1, rtp/broadinstitute.org/pub/ENCL_releases/release1/regional_misense_constraint/Sift 1 ensemble, released Jan 2013 <http://provean.jcvi.org/index.php>Sift 1402, released Nov 1, 2016 http://sift.bii.a-star.edu.sg/sift4public/Homo_sapiens/LRT 1, released November, 2009 http://www.genetics.wustl.edu/lrt/lrt_query.htmlMutation taster 2, data retrieved in 2015 <http://www.mutationtaster.org>MutationAssessor release 3, <http://mutationassessor.org/>PKUVEAN V2.1 ensemble, released Jan 2013 <http://provean.jcvi.org/index.php>FA1HMMP V2.3, <http://ratmm.biocompute.org.uk/ratmm-fa1hmmp/>ratmm-MKL, <http://ratmm.biocompute.org.uk/ratmm-mkl.htm>ratmm-X, <http://ratmm.biocompute.org.uk/ratmm-x/>

MetasVM and MetaLR, doi: 10.1136/annrheumdis-2013-204703

PrimateAI, <https://github.com/Illumina/PrimateAI>DeUGEN2, <https://deogen2.mutarrame.com/>ALoF 1.0, <http://aloft.gersteinlab.org/>BayesDel v1, <http://fengb-laboratory.org/BayesDel/BayesDel.html>LIST-S2 release: 2019_10, <https://precomputed.list-s2.msi.ubc.ca/>