SUPPLEMENTARY MATERIAL

SUPPLEMENTARY TABLE 1:

Reference	Study type	Populatio n/ Country	Objective	Sample Size	Source	Genes	SNP/Mutation	Results/conclusion
Adappa et al. 2016 [109]	CG	USA	To determine whether TAS2R38 genetics predicts outcomes in CRS patients following sinus surgery	82 CRSwNP 41 CRSsNP	NP, sinona sal tissue	TAS2R38	rs713598 (G/C; Ala/Pro)) rs1726866 (G/A; Val-Ala) rs10246939 (T/C; Ile-Val)	The genotype PAV/PAV was related to lower incidence of failing therapy and less frequent sinus surgeries
Ahmed et al. 2017 [33]	CG	Iraq	To clarify the role of <i>IL4</i> polymorphism in NP	22 healthy controls (HC) 36 NP	NP, inferio r turbin ate mucos a (ITM)	IL4	?	The polymorphism was found in NP patients but not in controls
Akygit et al. 2017 [53]	CG	Turkey	To identify genetic polymorphism of <i>SOD2</i> , <i>CAT</i> , <i>iNOS</i> enzymes in E- CRSwNP and NE- CRSwNP patients.	188 HC 65 E-CRSWNP 65 NE-CRSWNP	Blood	NOS2 SOD2 CAT	-277A/G 16C/T -21A/T	The GG genotype (NOS2) and TT genotype (CAT) distributions were different between E-CRSwNP and controls
Alromaih et al. 2013 [27]	pGWA S	Canada	To identify whether genetic factors associated with MHC1 deficiency are present in CRS	196 HC 154 CRSwNP 52 CRSsNP	Blood	CD8A TAPBP	rs3810831 rs2282851	The minor allele C in <i>CD8A</i> (OR 0.706; p=0.047) and heterozygous CT (OR 0.370; p=0.012) had a protective effect on the development of CRS. The minor allele T in <i>TAPBP</i> (OR 1.53; p=0.009) and heterozygous TT (OR 2.67; p=0.042) were associated with an increased risk for CRS.
Al-Shemari et al. 2008 [56]	CG	Canada	To evaluate the effects of SNPs on CRS in a panel of genes related to cysteinyl leukotriene metabolism	200 HC 179 CRSwNP 27 CRSsNP	Blood	ALOX5 AP	rs12430915 rs9506352 rs4769870 rs9579648 rs4076128 rs9579649 rs11616333 rs9315051 rs4769055 rs4420371 rs9578196 rs4466940 rs4293222 rs9578200 rs12429692 rs9285076 rs10162089 rs9670198 rs4254165 rs4319601 rs4356336 rs4769063 rs17612127 rs4238139	Three SNPs located within the ALOX5 (rs3780894), CYSLTR1 (rs321090) and ALOX5AP (rs17612127) genes reached the nominal p-value threshold (p < 0.05) for association with CRS. However, none of these SNPs resisted multiple testing adjustment.
							rs3780894 rs3780901 rs2279435 rs7099684 rs1565096 rs1487562 rs7919239 rs2291427 rs7393696 rs2115819 rs11239523 rs4948672 rs7089063	
						CYSLTR2	rs2406939 rs9595961 rs11617224 rs17072059 rs6420296 rs7330127 rs2407249 rs7335898 rs9568087 rs9285169 rs12184704	
						LTC4S	rs321090 rs321007 rs321006 rs730012	
							rs2291418 rs166624	
Bae et al. 2013 [110]	CG	Korea	I o investigate the association between <i>CIITA</i> and NP	All asthma patients: 158 CRSwNP 309 CRSsNP	Blood	CIITA	rs12932187 rs4781019 rs6498126 rs4781011 rs11074938 rs11074934 rs11074939 rs8043545 rs7404786 rs7201430 rs6498119 rs4781024 rs1139564 rs7189406 rs4774 rs6498124 rs4781016	I wo SNPs (rs12932187 and rs11074938) and 2 haplotypes (<i>CIITA</i> _BL1_ht2 and <i>CIITA</i> _BL1_ht5) were demonstrated to be associated with nasal polyps (P=0.001-0.01, OR=0.53-2.35 depending on the genetic model). After multiple testing correction only rs12932187 retained the association with nasal polyps (Pcorr=0.02).

Baldan et al. 2015 [111]	CG	Italy	To investigate the effect of 3 <i>IFRD1</i> SPNs on the development of NP in CF patients	CF patients: 40 with NP 103 without NP	Blood	IFRD1	rs6968084 (C>T) rs3807213 (A>C) rs7817 (C>T)	rs7817-CT showed 4-fold higher probability of NP than CC; the TT showed 7.3-fold increased probability. The CAT haplotype showed higher probability of NP (OR 2.63, p=0.004) compared to the CCC haplotype.
Batikhan et al. 2010 [47]	CG	Turkey	To investigate the possible association of <i>TNF</i> -308G/A with NP	95 HC 97 NP patients	Blood	TNF	rs1800629	The presence of the <i>TNF</i> -308 G/A SNP was an independent risk factor for development of NP (OR, 3.68; Cl, 1.27–10.7; p = 0.016)
Benito- Pescador et al. 2012 [57]	CG	Spain	To analyze polymorphisms in <i>LTC45</i> , <i>CYSLTR1</i> , <i>PTGDR</i> , and <i>NOS2</i> as representative genes of inflammation pathways in a population of patients with NP	245 HC 241 NP: 145 asthma 81 NSAID 75 aspirin triad	Blood	LTC4S CYSLTR1 PTGDR NOS2	rs730012 (-444A>C) 927 T>C -613 C>T -549 T>C -441 C>T -197 T>C CCTTT	The –444A>C <i>LTC4S</i> polymorphism was significantly associated with NP and atopy (P=.033) and with NP and atopic asthma, (P=.012). A significant association was found when the (CCTTT) repetition of the <i>NOS2A</i> gene was present more than 14 times in patients with NP and asthma (P=.034), in patients with P olyposis and intolerance to nonsteroidal anti-inflammatory drugs (P=.009), and in patients with the aspirin triad (P=.005). The <i>PTGDR</i> diplotype CCCT/CCCC (~513C, ~549CC, ~441CC and ~197TC) was more frequent in patients with NP (P=.043), NP with asthma (P=.013), and the aspirin triad (P=.041)
Berghea et al. 2014 [51]	CG	Romania	To investigate the association between <i>TNF</i> SNP with NP in Romanian patients with asthma	45 NP (38 NSAID+ 7 ATA) 61 without NP (8 NSAID+ 53 ATA)	Blood	TNF	rs1799724 (-857 C>T) rs1800629 (-308 G>A) rs361525 (-238 G>A)	There was an association of -857C>T with NP (p=0.01 ATA; p=0.05 NSAID). The allele T was more frequent in NP patients than in non-NP patients.
Bernstein et al. 2009 [30]	CG	USA	To investigate the role of 7 proinflammatory, 4 anti-inflammatory, one Toll receptor and 2 chemokine polymorphism in patients with massive NP	153 HC 179 NP	Buccal cells	TNF IL1A IL1B IL6 TGFB1 IL10 CD14 CCL5/ RANTES CCL2	rs1800629 rs3783521 rs17561 rs3087258 rs1143634 rs13447445 rs11466315 rs1800895 rs1800894 rs1800896 rs2569190 rs2107538 rs3917882	The frequency of the A allele in <i>TNF</i> is significantly higher in patients with NP versus controls (OR 1.86; 95% Cl, 1.4– 3.09)
Bohman et al. 2017 [21]	GWAS	Sweden	To identify genetic markers and genes associated with susceptibility to CRSwNP using a family- based GWAS	393 HC 427 CRSwNP	Blood	HLCS HLA-DRA BICD2 VSIR SLC5A1		Pathway analyses using top 1000 markers with the most significant association p- values resulted in 138 target genes. Comparisons with data from expression quantitative trait loci showed the most skewed allelic distributions in cases with CRSwNP compared with HC for the genes HLCS, HLA-DRA, BICD2, VSIR and SLC5A1
Bosse et al. 2009 [73]	GWAS	Canada	To perform pooling- based GWAS in two case-control cohorts, one of them consisting of patients with CRSwNP	189 HC 210 with severe CRS 157 CRSwNP 53 CRSsNP	Blood	LAMA2 PARS2 NAV3 LAMB1	rs2571584 rs2873551 rs1726427 rs4727695	600 SNPs from 445 genes that were potentially associated with CRS (P < 0.05). Each of these novel high-priority SNPs had allele frequency differences between cases and controls at a level worthy of additional investigation. The most significant SNP for each of the top 10 genes are shown in this table.

-		r	1	I		1	1		
							CACNA1I	rs3788568	
							KIAA1456	rs11779957	
							MUSK	rs10817091	
							TRIP12	rs10535833	
							AOAH	rs4504543	
							MSRA	rs7001821	
Bussu et al. 2007 [69]		CG	Italy	To evaluate the potential involvement of ADRB2 A16G polymorphism in sinonasal polyposis	47 HC 56 NP	Blood	ADRB2	rs1042713 (g.5285A>G)	The presence of Arg (A allele) is significantly higher in NP patients than in controls (p=0.0386)
Buysschaert		GWAS	Belgium	To investigate whether	415 HC	Blood	IL1RL1	rs1420101	Rs3939286 was significantly associated
et al. 2010 [41]				certain SNPs predispose to NP	273 NP		IKZF2	rs12619285	with NP (OR 1.60; 95% CI 1.16-2.22; p=0.0041). The A-allele conferred a risk for NP (OR 1.53; 95% CI 1.21-1.96;
							GATA2	rs4431128	p=0.0041).
							IL5	rs4143832	Rs1420101 may increase risk when in combination with rs3939286
							SH2B3	rs3184504	
							WDR36	rs2416257	
							мнс	rs2269426	
							МҮВ	rs9494145	
							GFRA2	rs748065	
							IL33	rs3939286	
Cantone et al. 2018 [61]		CG	Italy	To investigate the relevance of TAS2R38	100 CRSwNP	Saliva, blood	TAS2R38	rs713598 (C145G; Pro>Ala)	The nonfunctional genotype (AVI) is more frequent among CRS patients than in the
				genetic variants in the susceptibility to				rs1726866 (C785T; Ala>Val)	general population (25% vs. 18.4%, P=0.034). No relationship with severity
				bacterial infections				rs10246939 (G886A;Val>IIe)	was found.
Castano et al. 2009 [36]		CG	Canada	To investigate whether whether certain	187 HC	Blood	IL1RL1	rs974389 rs10204137 rs985523 rs10208293	Statistically significant allelic associations with CRS were noted for 5 SNPs
				polymorphisms in the	154 CRSwNP			rs1041973 rs12105808	(rs10204137, p=0.04; rs10208293, p=0.03;
				IL1RL1 gene are differentially present in	52 CRSsNP			rs1420103 rs12712142	rs13431828, p=0.008; rs2160203, p=0.03,
				patients with surgery- unresponsive CRS and in				rs1921622 rs12996097 rs2160203 rs13431828	and rs4988957, p=0.03).
				control subjects				ro2771177 ro17606274	But only one SNP significantly associated with CRSwNP (rs13431828, n=0,008)
								rs4988957	with crown (1919491020, p=0.000)
Castano et		CG	Canada	To assess the	196 HC	Blood	MET	rs38840 rs2237711 rs1024658	The genotype distribution of two SNPs in
al. 2010 [71]				association of polymorphisms in the	154 CRSwNP			rs10271561 rs10243024	the <i>MET</i> gene
				MET gene with CRS in a Canadian population	52 CRSsNP			rs40239 rs714180 rs38855	(rs388840, rs38850) displayed a significantly different genotypic
								rs38841 rs38857	distribution (p≤ 0.05) between CRS
								rs39747	patients and controls. The most significant association in the MET gene
								rs38845 rs38846 rs2200440 rs1752600	was found with SNP rs38850 (p=0.004).
								rs7798983 rs2402118	
								rs38849	
								rs722134 rs38850 rc1621	
								131021	
Chang at al		66	Taiwan	To assess the	103 HC	Blood	II 1P	1542530	There were significant differences in the
2006 [34]			i aiw/d[]	association of <i>IL1B</i> and	88 CPC	воод	ILID	+3953C5T	distribution
				polymorphisms with CRS	61 CRSwNP		IL1RN	Variable number tandem repeat of	of the IL1RN polymorphism between the control
					27 CRSsNP			an 86–base pair segment in intron 2	subjects and patients with CRS (P=.05).
									of <i>IL1RN</i> occurred more frequently in the
									CRS patient

								group (OR 3.3; 95% CI, 1.25-9.18, P=0.01).
Cormier et al. 2009 [112]	CG	Canada	To determine whether SNP in <i>TNF, TNFAIP3</i> , and <i>TNFAIP6</i> genes were associated with CRS	196 HC 206 CRS 154 CRSwNP	Blood	TNF TNFAIP3 TNFAIP6	rs2229094 rs3093672 rs1121800 rs769177 rs1321136r rs77669888 s1800750 rs9267502 rs2256965 rs9469027 rs2256974 rs1800629 rs2857706 rs361525 rs3093561 rs4987027 rs5029935 rs5029938 rs5029939 rs661561 rs610604 rs50 29965 rs6433371 rs16830015 rs670782 rs10183099 rs2342910 rs3771891	Two polymorphisms in <i>TNFAIP3</i> (rs3757173 and rs5029938) are weakly associated with severe CR5 but no association was found with genetic variants in TNF or <i>TNFAIP6</i> . None was associated to risk of NP.
Dar et al. 2018 [65]	CG	India	To assess the risk of CR5wNP conferred by SNPs in <i>FccR1α</i> gene in a North Indian cohort	50 HC 100 CRSwNP	Blood	FCER1A	rs10432475 rs2427827 rs2251746 rs2298804 rs2298805 rs2269718	In those cases with high serum IgE, the T allele of rs2427827 is significantly more frequent in CRSwNP patients (OR 2.24; p=0.02)
De Alarcon et al. 2016 [58]	CG	USA	To evaluate the association of <i>LTC4S and</i> <i>PAI-1</i> variants with CRS	66 HC 16 CRSsNP 117 CRSwNP	Blood, polyp fibrobl asts	LTC4S SERPINE1 (PAI1)	rs730012 (-444A>C) rs1799762 (4G/5G ins.)	The C allele of <i>LTC4S</i> was more frequent in those NP patients also diagnosed with chronic hyperplastic eosinophilic sinusitis (p<0.04)
Ekinci et al. 2011 [70]	CG	Turkey	To examine whether there is an association of eotaxin-1 (<i>CCL11</i>) gene polymorphisms with NP	93 HC 85 NP	Blood	CCL11	rs1490392522 (-384A>G) rs762429865 (67 G>A)	The selected SNPs are more frequent in NP patients than in HC (p=0.044 and p=0.019, respectively). However, their relation was statistically poor (association coefficient =0.18).
Erbek et al. 2007 [35]	CG	Turkey	To investigate the association between NP and SMPs of the proinflammatory cytokines <i>IL1A</i> , <i>IL1B</i> , <i>TNFα</i> .	106 HC 82 NP	Blood	IL1A IL1B TNF	4845G>T -511C>T rs361525 (-238 G>A) rs 1800629 (-308G>A)	The 4845GTand 4845TTgenotypes of the <i>IL1A</i> gene were associated with NP (P<.05). The frequency of the <i>IL1B</i> –511 CC and CT were significantly higher in patients with NP than in controls (P=.01). There was a significantly high risk of susceptibility to NP in patients with the -308 GA genotype (P=.001).
Esmaeilzade h et al. 2015 [19]	CG	Iran	To investigate the association of <i>HLA-DRB</i> and - <i>DQ</i> genetic variabilities in patients with AERD	100 HC 50 CRSwNP + asthma	Blood	HLA- DRB1	HLA-DRB1*0101 HLA-DRB1*15 HLA-DRB1*16 HLA-DRB1*0301 HLA-DRB1*04 HLA-DRB1*07 HLA-DRB1*08 HLA-DRB1*0901 HLA-DRB1*1001 HLA-DRB1*110 HLA-DRB1*110 HLA-DRB1*1301 HLA-DRB1*1302	Two variations are associated with increased risk of NP: <i>HLA-DRB4</i> , OR 2.34, CI 1.37–4.00, p>0.01 <i>HLA-DQB1*0302</i> , OR 4.56, CI 2.10–9.91, p<0.01 Two variations are associated with reduced risk of NP: <i>HLA-DRB3</i> , OR 0.41, CI 0.24–0.68, p<0.01 <i>HLA-DQB1*0301</i> , OR 0.39, CI 0.21–0.73, p<0.01

-									[
								HLA-DRB1 1305	
								HLA-DRB1*14	
							HLA- DRB3	HLA-DRB3	
							HLA-	HLA-DRB4	
							DRB4		
							HLA- DRB5	HLA-DRB5	
							0.000		
							HLA- DQA1	HLA-DQA1*0101	
								HLA-DQA1*0102	
								HLA-DQA1*0103	
								HLA-DQA1*0104	
								HLA-DQA1*0201	
								HLA-DQA1*0301	
								HLA-DQA1*0401	
								HLA-DQA1*0501	
							HLA-	HLA-DQB1*0201	
							DQB1	HLA-DQB1*0301	
								HLA-DQB1*0302	
								HLA-DOB1*0303	
								HLA-DOB1*0305	
								HLA-DOB1*0401	
								HIA-DOB1*0501	
								HLA-DOB1*0601	
								HLA DOB1*0602	
								HLA-DQB1 0002	
								HLA-DQB1*0604	
Fajardo- Dolci et al.		CG	Mexico	contribution of the	151 HC	Blood	HLA- DQA1	HLA-DQA1*0101/4	Ine allele <i>HLA-DQA1*0201</i> was found to be involved in susceptibility to develop
2006 [20]				human major histocompatibility	31 NP			HLA-DQA1*0102	simple NP, without asthma, aspirin intolerance, or any other allergic
				complex HLA-DQA1, – DQB1, and TNF α genes				HLA-DQA1*0103	diseases. OR 6.79 CI (1.9-23.9)
				with simple nasal polyposis.				HLA-DQA1*0201	13% etiological fraction was found for the haplotype HLA-DQA1*0201-DQB1*0201
								HLA-DQA1*030101	(P=0.016).
								HLA-DQA1*0401	
								HLA-DQA1*0501	
	-						HLA-	HLA-DQB1*0201	
							DQBI	HLA-DQB1*0301	
								HLA-DQB1*0302	
								HLA-DQB1*0402	
								HLA-DQB1*0501	
								HLA-DQB1*0502	
								HLA-DOB1*0503	
								HIA-DOB1*0504	
								HLA-DQB1*0601	

					r	1	HLA-DOB1*0602	
							HLA-DOR1*0603	
							TEA-DODI 0003	
						TNF	rs1800629 (-308 G>A)	
							rs361525 (-238 G>A)	
Fruth et al. 2011 [113]	CG	Germany	To analyze the potential association of GST	52 HC	NP, ITM	GST	GST-T1	No correlation
			polymorphisms and CRS.	118 CRS			GST-M1	
				69 CRSwNP			GST-P1	
				49 CRSsNP				
Fruth et al	60	Germany	To shed light on the	30 HC	NP	SPINK5	rs17775319 (G1258A)	No correlation
2012 [114]		Germany	significance of SPINK5	50 000000	ITM	51 1110	C2475T	
			inflammatory diseases	59 CRSWINP			624751	
			of the upper respiratory tract.	15 CRSsNP			rs1243172589 (A2915G)	
							rs745601984 (A1103G)	
Gallo et al.	CG	Italy	To confirm the	39 HC	Blood	TAS2R38	rs713598 (G/C; Ala/Pro))	No differences found in genotypic
2010 [02]			between TAS2R38	36 CRSwNP			rs1726866 (G/A; Val-Ala)	distribution
			genotype, CRS, and related comorbidities.	17 CRSsNP			rs10246939 (T/C; Ile-Val)	
Henmyr et	GWAS	Turkey.	To investigate the	1588 HC from	Blood.	II.1A	rs17561	Some SNPs are associated with increased
al. 2014 [37]	0	Finland,	reproducibility of	Illumina data	databa	,		risk of NP:
		Korea, USA,	associations with	base	se			IL1A rs17561
		Belgium	CRSSNP and CRSWNP.	613 Belgian patients:		IL1B	rs16944	<i>RYBP</i> rs4532099
				275 CRSwNP		RYBP	rs4532099	TNF rs1800629
				338 CRScND		DCBLD2	rs828618	// 22 **202028C
	-			556 CH55INF		TNFA	rs1800629	1633 153339280
							rs361525	<i>IL10</i> rs1800870
	-					10111		CACNG6 rs192808
						AUAH	rs4504543	MMP9 rs3918242
						IL33	rs3939286	
						IRAK4	rs4251431	Some SNPs are associated with reduced
	-					IL10	rs1800870	risk of NP
						CACNG6	rs192808	<i>IL1B</i> rs16944
						MMP9	rs3918242	DCBLD2 rs828618
							rc17577	AOAH rs4504543
							121/2//	IRAK4 rs4251431
Henmvr et	CG	Sweden.	To screen for rare	372 HC	Blood	PARS2	rs143717155 rs116816976	A significant surplus of variation was
al. 2016		1000Genom	variants in PARS2 and to	120 CDCurbID			rs35201073 rs12023572	observed in the CRS patients (p=0.048).
[113]		con oject	accumulation of such	120 CK2WNP			rs370234936 rs61768813	Haplotype analysis of the region showed
			variants in CKS patients.	172 CRSsNP			rs2270004 rs145005088 rs563439229	a significant excess of rare haplotypes in the CRS patients compared to HC in the
							rs1180946 rs1180945 rs11577368 rs116416055	following SNPs:
							rs145866387	rs2873551, rs2270004, rs11577368, rs1180946, rs1180945
							rs74617964 rs1180947	TTAGC n=0.0049
								11CCC p=0.0048
								TCAGT p=0.0016
Hytönen et	CG	Finland	To investigate if the frequency of the most	135 CRS	Blood	CFTR	ΔF508	No abnormal distribution was observed in CER patients
ai. 2001 [07]			common CFTR					Ciri patienta
			mutations was more			1		

			common among CRS patients with or without NP.	91 CRSsNP 46 CRSwNP			394delTT	
Ismi et al. 2017 [50]	CG	Turkey	To determine the genetic susceptibility of NP formation to TNF	91 HC 71 CRSwNP	Blood	TNF	-308G>A	There was a statistically significant increase in the expression of the <i>TNF</i> -308 GG and <i>IL1B</i> -511 CC genotypes in the
			and IL1B polymorphisms			IL1B	-511T>C	patients with NP
Karjalainen et al. 2003	CG	Finland	To establish whether IL1A and IL1B have an	35 CRSwNP	Blood	IL1A	4845G>T	The risk of NP was markedly increased in IL1A allele G homozygous subjects (OR
[38]			effect on susceptibility to NP.	210 CRSsNP		IL1B	-511C>T	2.73; 95%CI 1.40–5.32, p=0.005). In the case of <i>IL1B</i> no significant associations were found.
Keles et al.	CG	Turkey	To evaluate whether	100 HC	Blood	HLA-A	HLA-A *01	HLA-B*07 and -Cw*12 alleles were
2008 [22]			there is a relationship between HLA-A, -B, -Cw, and -DRB1 alleles and	66 NP			HLA-A *02	than in the control group.
			developing NP.				HLA-A *03 HLA-A *11	HLA-B*57 and -Cw*04 alleles were significantly lower in the NP patients than in the control group.
							HLA-A *24	In the NP patients with ASA, there was a
							HLA-A *26	HLA-A*24, -Cw*01, -Cw*12, and - DRB1*04 alleles.
							HLA-A *33	HLA-A*33 and -Cw*12 alleles in NP
						HLA-B	HLA-B *07	patients who had polypectomy history were significantly higher than in the control group
							HLA-B *15	In NP patients, a significantly decreased
							HLA-B *35	frequency of the HLACw*
							HLA-B *51	04 and -DRB1*11 alleles was shown.
							HLA-B *57	
						HLA-Cw	HLA-Cw *01	-
							HLA-Cw *02	
							HLA-Cw *03	
							HLA-Cw *04	
							HLA-Cw *06	
							HLA-Cw *07	
							HLA-Cw *08	
							HLA-Cw *12	-
						HLA- DRB1	HLA-DRB1*01	
							HLA-DRB1*04	
							HLA-DRB1*07	
							HLA-DRB1*11	
							HLA-DRB1*14	
Kilty et al. 2010 [116]	CG	Canada	To investigate the association between	196 HC	Blood	SERPINA1	rs11558262 rs1956707 rs11832	Two SNPs (rs1243168 and rs4900229) were associated with the disease.
			SNPs in the SERPINA1 gene and CRS	154 CRSwNP			rs2071274 rs1243160	rs1243168 T allele was significantly
				27 CRSsNP			rs1243163 rs2230075	associated with severity (p<0.01)
							rs1243166 rs2239651 rs1243167	
							rs1243168 rs1243169 rs2753934	
							rs1243171 rs3748316 rs12884390	
							rs4900229 rs1303 rs17287271 rs4900230	
							rs17580	

							rs17751614	rs4905198	
							rs17751769 rs17824797	rs6575424	
								rs6647	
								rs709932	
								rs7151526	
								rs8010121	
								rs877081	
Kim et al.	CG	Korea	To evaluate the	456 HC	Blood	TGFB1	rs13447445		No association with NP
2007 [117]			association of TGFB1 polymorphisms with an	206 AERD					
			AERD phenotype in the Korean population	72 NP					
				324 ATA					
				10 NP					
Kim at al		Karaa	To evoluete esseciations	192.110	Disad	404	m11086033		No association with ND was described
2009 [64]	6	Korea	between genetic	136 AERD	ыооч	ADA	1511080932		Cignificant differences between normal
			adenosine deaminase and adenosine	130 AERD		400841	15244076		and patients with
			receptors with the AERD phenotype	47 NF 181 ATA		ADURAI	rs6664108		AERD in the ADORA1 SNP genotype frequencies for rs16851030 (P=0.001) and
				10 NP			rs6427994		rs6664108 (P=0.013).
							rs16851030		
							rs12744240		
						ADORA2A	rs5996696		
							rs5751876		
						ADORA3	rs2298191		
							rs10776727		
							rs1544224		
							rs2229155		
Kim et al. 2012 [23]	CG	Korea	To investigate the associations of HLA-DRA	158 CRSwNP	Blood	HLA-DRA	rs9268628 A>0	C	4 SNPs were significantly associated with NP
()			polymorphisms with NP in asthmatic and AERD	309 CRSsNP			rs3129871 C>/	Ą	Rs9268644 p=0.009
			patients.				rs9268633 G>	4	Rc3129878 n=0 033
							rs9405035 G>/	Ą	D-2120881 p=0.012
							rs14004 C>A		RS3129881 p=0.013
							rs9268644 C>/	4	Rs2239805 p=0.029
							rs9268645 G>	с	And the haplotype (rs3129871; rs8084; rs2239805; rs2239804; rs7192;
							rs3129878 A>0	C	rs4935354; rs7194; rs1051336; rs1041885) TAAATGGA (p=0.029)
							rs3135392 G>	г	
							rs6926374 G>/	A	
							rs3129881 C>	г	
							rs17496549 C	۶T	
							rs6911777 T>0	2	
							rs3129886 C \1	-	
							rs8084 (>4		
							rs222080E A	~	
							132237003 A>	~	
							152239804 G>	•	
	1		1				rs/192 G>T		
							rs4935354 A>(3	

							rs1051336 G>A	
							rs1041885 A>T	
Kostuch et al. 2005 [66]	CG	Poland	To determine the prevalence of the most common CFTR mutations in patients with NP without suspicion of CF.	70 human placentas 44 NP	Blood, placen ta	CFTR	ΔF508 G551D G542X N1303K	ΔF508 is more frequent in patients than in HC (p=0.0032) and in the general Polish population as well (P =0.0059).
							1717-1G>A W1282X R553X ΔΙ507	
Kosugi et al. 2013 [118]	CG	Brazil	To analyze the relationship between an IL6 polymorphism and asthmatic NP patients.	81 HC 45 asthmatic with NP 63 non asthmatic NP 45 asthmatic without NP	Blood	IL6	rs374295772 (-174G>C)	Genotype distribution was non- significant, but GG genotype appeared more frequently in all inflammatory groups.
Kristjansson et al. 2019 [55]	GWAS	Iceland, UK	To search for sequence variants affecting the risk of NP or CRS	Iceland 353939 HC	Databa se	HLA- DQA1	rs1391371	The mentioned variants at ten loci were found that associate with NP at genome- wide significance.
				3188 cases		IL33	rs1888909	The variant with the largest effects on the
				UK		TSLP	rs1837253	risk of NP is a low-frequency missense variant rs34210653[A] (Thr560Met) in
				406147 HC		ALOX15	rs34210653	ALOX15 that confers a 68% reduction in NP risk (p= 8.0x10 ⁻²⁷
				2420 cases		10p14	rs1444782	OR 0.32, 95% CI 0.26, 0.39).
				2420 00303		FOXP1	rs17718444	
						CYP2S1	rs338598	
						IL18R1	rs6543124	
						SI C22A4	rs1050152	
						MYRF	rs174535	
Kuran et al.	CG	Turkey	To analyze possible	98 HC	Blood	IL1RN	rs2234663	Distribution of genotypes of IL1RN and
2015 [55]			increase susceptibility	78 NP		IL4	rs8179190	(p=0.0001)
			to Nr.			IL2	rs206976	
Luxenberger et al. 2000	CG	Austria	To determine the association of HLA-A, -B,	1070 HC	Blood	HLA-A		A significant association was seen with HLA-A74
[119]			-DR, and –DQ with NP	89 NP		HLA-B		and nasal polynosis
						HLA-DR		
						HLA-DQ		
Mfuna Endam et al. 2009 [29]	CG	Canada	To explore association between SMPs in IL22RA1 and severe CRS	196 HC 206 CRS 154 CRSwNP 52 CRSsNP	Blood	IL22RA1	rs10465895 rs2502450 rs10751768 rs3795300 rs10794665 rs10903031 rs3936073 rs11249201 rs4292900 rs11577442 rs4648936 rs11578307 rs11578657 rs4648942 rs12070496 rs460187	Three SNPs (rs4292900 Pnom =0.0006, OR 1.757; rs4648936 Pnom=0.0011, OR 1.716; rs16829225 Pnom=0.0014, OR 1.977) show significant differences in allelic frequencies between cases and controls
Mfuna Endam et al. 2010 [28]	CG	Canada	To replicate the CRS associations recorded	196 HC	Blood	ILIA	rs12092673 rs6424157 rs12408946 rs16829225 rs7418238 rs7513249 rs17561 rs1800587 rs783521	For rs17561, this study replicated previous results about the association of the TT homozygote genotype (OR, 3.39;
			for IL1A, IL1B, and TNF				rs2048874	P=.007). The protective effect of

				in a cohort of Canadian	206 CRS			rs2856838	rs6722023	rs2856838 (OR, 0.38; P=.002) and the risk
				patients with severe CRS.	154 CRSwNP		IL1B	rs16944		effect of rs1800587 (OR=3.16, P=.008) are enhanced with the homozygote form of the minor allele
					52 CRSsNP		TNF	rs1121800	rs3093561	
								rs13211368	rs3093672	with SNPs in IL1B or TNF.
								rs1800629 rs1800750	rs361525	
								rs2229094	rs4987027	
								rs2256965	rs769177	
								rs2256974 rs2857706	rs7766988	
									rs9267502	
									rs9469027	
Mfuna		GWAS	Canada	To identify taste	GCRS1	Blood	TAS2R1	rs17788846	rs6874254	57 SNPs in TAS2Rs and 16 SNPs in TAS1Rs
Endam et al. 2014 [59]				receptor associated with CRS and verify	196 HC			rs41483	rs882142	had allele frequency differences above 10% between controls and patients
				whether known SNP replicated in their CRS	206 CRS			rs12374524 rs6555620	rs4272105 rs11739710	(range, 10.2% to 32.4%).
				cohort				rs10746553	rs3110986	Three coding SNPs associated with CRS were identified: 1 in the TAS2R13 gene
					GCR52			rs1015855	rs3094363	(rs1015443, biallelic differences of 13.8% in GCRS1), and 2 others in the TAS2R49
					190 HC		TA\$2810	rs669503		gene (rs12226920, biallelic difference of 16.0% in GCRS1; and rs12226919, biallelic
					408 CRSwNP			rs10845219		difference of 11.9% in GCRS1)
							TAS2012	rc101E442		
							1432113	151015442		
							7462044	151015445		
							TAS2R14	rs3851586		
								rs1013311		
								rs3741843		
							TAS2R3	rs765007		
								rs6962760		
							TAS2R38	rs4726481		
								rs10246939		
								rs1726866		
							TAS2R39	rs11979433		
							TAS2R4	rs2234002		
								rs2190245		
							TAS2R41	rs2966709 rs2966715	rs12536735	
								rs2949746	rs1806578	
								rs2949770	rs2966701	
									rs2966699	
							TAS2R43	rs2966699	rs2708333 rs2597975	
								rc2708264	rc7313682	
								rs2599396	rs2597974	
							TAS2R44	rs4763616		
								rs2010481		
							TAS2R46	rs2708389		
								rs11533164		
								rs2708377		
								rs2255418		
							TAS2R48	rs10772420		
1	1	1	1	1	1	1	1	1		1

						TAS2R49	rs1463237 rs10772408 rs4298989 rs12226920	
							rs12581501 rs12226919 rs11054150	
						TAS2R5	rs35010424	
							rs11769672	
							rs11773137	
							rs1859646	
						TAS2R50	rs2900554	
							rs6488331	
						TAS2R7	rs7313019	
						TAS1R1	rs11122100	
							rs12080675	
						TAS1R2	rs28410948 rs6686865 rs7417542	
							rs9662598 rs7411042	
							rs4920566 rs6685177 rs1	2137730
							rs6603912 rs6684577 rs12063142	
							rs3935570 rs12042960 rs7418296	
Molga et al. 2016 [120]	CG	Poland	To assess genetic predisposition of	463 HC	Blood	MMP1	rs199750 (-1607 G/GG)	The frequency of genotypes was not significant related to CRSwNP. but GG is
()			MMP1 -1607 G/GG	206 CRSwNP				relates to increases number of surgeries (p=0.002) and bronchial asthma (p=0.021)
			polymorphism to CRSwNP					
Molnar- Gabor et al.	CG	Hungary	To investigate whether there is an association	50 HC	Blood	HLA-DR5		The odds ratios for developing nasal polyposis increased in people carrying the
2000 [24]			Detween HLA-DRB1, - DQA1, and DQB1 alleles	50 CRSwNP		HLA-DR7		DR7 (OR 2.55) allele with the linked DQA1*0201 (OR 2.52) and DQB1*0202
			and developing w			HLA- DQA1	HLA-DQA1*0101	DR5 (OR 0.66) linked with DQA1*05012 (OR 0.69) DOB1*0301 (OR 0.57) alleles
							HLA-DQA1*0102	showed a decreased odds
							HLA-DQA1*0103	ratio value.
							HLA-DQA1*0104	
							HLA-DQA1 0201	
							HIA-DOA1*05011	
							HLA-DQA1*05012	
						HLA-	1-10	
						DRB1		
						HLA- DQB1	HLA-DQB1*0201	
							HLA-DQB1*0202	
							HLA-DQB1*0302	
							HLA-DQB1*0301	
							HLA-DUB1*0303	
							HLA-DOB1*0502	
							HIA-DOB1*0601	
							HIA-DOB1*0602	
							HIA-DOB1*0603	
							HLA-DUDI 0003	

							HLA-DQB1*0604	
Mrowicka et	CG	Poland	To investigate the	200 HC	Blood	IL1B	rs55778004 (-511C>T)	The TT genotype for C-511T mutation
al. 2014 [32]			relationship between IL1B	208 CRSwNP				associated with
			and II 4 promoter				5000 T	the risk of developing NP in a Polish
			polymorphisms			11.4	-590C>1	population
Nakayama	GWAS	Japan	To perform an	1908 HC	Blood	TSLP	rs1751303 rs3806933	Significant association between CRSwNP
et al. 2020 [121]			association study of CRSwNP and AERD with	499 CRSwNP			rs10056340 rs1898671	and rs1837253, rs3806932 and rs3806933, with the most significant
			genetic variants in the TSLP locus				rs1837253 rs2416257	association being observed at rs1837253 (p= 1.27x10 ⁻⁶ ; OR, 1.60; 95% CI, 1.32-
							*******	1.94)
							153800932 151438073	
Ozdas et al. 2015 [122]	CG	Turkey	To analyze SNPs of the RYD5 gene and to	238 HC	Blood	RYD5	rs113795008 rs7951297	Four SNPs (rs113795008, rs2280540, rs2294083, and rs2294082)
			determine the effect on polyp formation	196 NP			rs535294582 rs2294083	were significantly associated with NP. The
							rs2280540 rs2294082	individuals with combined genotypes of six risk alleles (rs113795008 rs2280540
							rs144999256 rs61997072	rs7951297, rs2294083, and rs2294082)
							rs148962288	compared with the ones with one or four
								risk alleles.
Palikhe et al. 2017 [123]	CG	Korea	To investigate the potential associations	120HC	Blood	ABCC4	rs868853 (A>G)	No significant association
			between ABCC4 gene polymorphisms and	270 asthma			rs839951 (C>G)	
			asthma genotype.	106 NP				
Park et al.	CG	Korea	To evaluate expression	70 HC	Blood	IL4	-590 C>T	A T>C exchange at -590 position was
2006 [43]			of cyclooxygenase (COX)-2 and 5-	61 NP				correlated with NP. The T allele was significantly more frequent in NP
			lipoxygenase (5-LO) associated with IL4					(p=0.028).
			promoter polymorphism -590 in NP tissues					
Pasaio et al	66	Koroa	To ovaloro the	200 HC	Databa	DCRLD2	rc2420224 rc929619	Six SNDs were associated with the
2012a [124]	69	Kolea	association of DCBLD2	509 HC	se	DCBLDZ	152455224 15626016	presence of NP:
			of NP in asthma patients	158 NP+asthma			rs1371687 rs828616	rs1371687, rs7615856, rs828621,
							rs9838238 rs16840208	rs828618, rs828616, and rs8833. After multiple testing adjustment, only
							rs17278047 rs17270986	rs828621 remained significant (p=0.006)
							rs7615856 rs1062196	
							rs828621 rs8833	
Pasaje et al.	CG	Korea	To investigate the	309 asthma no	Databa	EMID	rs6945102 rs1476652	Ten EMID2 SNPs (rs6945102, rs4729697,
2012b [125]			association between EMID and NP	NP	se		rs4729697 rs6973489	rs221, rs10435333, rs6947185, rs4727494, rs13233066, rs1008064,
				158 asthma+NP			rc10227610 rc7902156	rs1543883, and rs13245946) were associated with the presence of pasal
							1510257010 157802150	polyps (p= 0.004- 0.05, OR 0.61-1.32)
							rs9986717 rs10953342	depending on the genetic model.
							rs10254516 rs12668018	rs6945102, rs4729697, rs221, and rs10435333, were found to be
							rs10239458 rs1008064	significantly associated with NP in the overall Korean asthma patients even after
							rs221 rs13232646	multiple testing corrections
							rs10435333 rs1543883	
							rs6944691 rs1859625	
							rs6942770 rs6947089	
							rs9640666 rs9969331	
							150947185 IS12538381	
							rs11770876 rs17135512	
							rs11772022 rs1558015	
							rs11772003 rs10250055	
							rs10223928 rs6947735	
							rs4729705 rs2158739	
							rs10254310 rs10279545	
	1				1			

							rs4045	rs6945961	
							rc60/0700	rc13245946	
							rs4727491	rs17470799	
							rs13238748	rs10237510	
							rs4727494	rs17135617	
							m12222066	rc17125621	
							1513233000	1517155621	
							rs869127		
Pascual et al. 2008	CG	Spain	To analyze the (CCTTT)n polymorphism of NOS2	98 HC	Blood	NOS2	(CCTTT)n		Allele frequency distribution is significantly different between HC and NP
[126]			and/or asthma	46 NP					associated with increased risk of NP (OR
				150 asthma					14.39, 93% CI, 3.02 - 68.60, P = .001)
Pavon- Romero et	SNP array	Mexico	To evaluate whether contribution to	179 HC	Blood	ACE	rs4309†		In AERD vs. HC, we identified 22 associated SNPs, with 11 SNPs
al. 2018 [54]			reported in other	120 AERD			rs4293†		associated with risk in 7 genes (ACE,
			associated with AERD in Mexican nations	179 asthma		MS4A2	rs576790†		ANX4; denoted as t in the adjacent
			inexical patients				rs502581†		associated: ACE rs4309 (C allele p = 0.0001 OR = 1.92 CL 95% = $1.37-2.69$)
						FSIP2	rs2631700†		and MS4A2 rs573790 (C allele p = 0.0002, OR = 1.94 CL95% = 1.35–2.79)
							rs2631702†		By contrast 11 SNPs in 5 genes (PPARG
						KIFC3	rs2285700†		IL10, RG7SBP, TBXAS1, and FANCC) were associated with protection.
						ANX4	rs7588022†		·····
						FCER1G	rs4489574†		
							rs7528588		
						IL10	rs1800896†		
							rs3024498†		
							rs1554286		
							rs1800872		
						PPARG	rs2960421		
							rs4135275		
							rs1875796		
						RGS7BP	rs6870654		
						TBXAS1	rs13239058		
							rs10487667		
							rs6962291		
						FANCC	rs1326188		
Purnell et al. 2019 [63]	CG	USA	To determine the frequency of 6 SNPs in	1000 Genomes database	Buccal cells	TAS2R38	rs713598		No differences between CRSwNP and CRSsNP
[00]			genes with bitter taste signaling function.	74 CRS		GNB3	rs5443		
			-9	41 CRSwNP		TAS2R19	rs10772420		
				33 CRSsNP		TAS2R20	rs12226920		
						RGS21	rs7528947		
							rs1175152		
Ramirez-	CG	Mexico	To determine the	99 HC	Blood	HLA-	HLA-DRB1*02		Significant increase in the *03 and *04
al. 2006 [25]			association of HLA-DRB1 alleles with NP in the Mexican Mastiza	34 NP		DKB1	HLA-DRB1*03		(UK 2.2, p=0.009) allele trequencies.
			population.				HLA-DRB1*04		Significant decrease in the *08 allele (OR 0.2, p=0.01)
							HLA-DRB1*05		
							HLA-DRB1*07		
							HLA-DRB1*08		

Sachse et al. 2010 [127]	CG	Germany	To detect staphylococcal colonization in nasal	51 HC 68 NP	NP, ITM	TLR2	rs5743708	The minor allele A is not associated with NP
Sitarek et al. 2012 [52]	CG	Poland	To investigate the	200 HC	Blood	COX2	rs20417	Increased risk (p>0.001) of CRSwNP
2012 [52]			and MET gene polymorphisms with the risk of CRSwNP.	195 NP		MET	rs78116323	6.05) and G allele of MET (OR 5.52) The combined genotype GC/GG had
								increased risk (OR 4.07, p<0.001)
Song et al. 2012 [128]	CG	Korea	To investigate the genetic contribution of ALO15 to the development of AERD.	195 HC 171 AERD (49 NP)	Blood	ALOX15	rs34104097 rs7220870 rs2664592	The patients carrying haplotype 1 (GCG) of Rs34104097, Rs7220870, and Rs2664592 showed a significantly higher total eosinophil count compared to the other haplotypes (p = 0.008) in the AERD
				229 ATA (9 NP)		74/5	4000520	group
2013 [49]		nungary	TNFa -308G>A SMP has a role in the genetic predisposition to CRS in a Hungarian population.	326 CRSwNP 49 CRSsNP	cells	INF	12190053	A allele-containing genotypes among the AIA CRSwNP patients
Szabo et al.	CG	Hungary	To examine whether the	169 HC	Buccal	TNF	rs1800629	Carriers of 8.1 AH carried all 4 studied
2015 [48]			association between TNFa -308A allele and AIA CRSwNP is due to	244 CRSwNP	cells	AGER	rs1800625	SNPs in homozygotic or heterozygotic forms. This AH is significantly associated with CRSwNP (p=0.014)
			this allele or to the presence of the	57 CRSsNP		HSP70-2	rs1061581	-
			haplogroup (AH) in chromosome 6.			LTA	rs909253	-
Tewfik et al.	CG	USA	To investigate whether	154 CRSwNP	Blood	TLR11	rs4286521 rs4833095	Blood IgE levels have been shown to be
2009 [31]			polymorphisms in the genes encoding key TLR	27 CRSsNP			rs5743611 rs5743594	raised in patients with CRSwNP
			signaling molecules might be associated				rs4833103	The C allele of rs1461567, the G allele of rs4251513, and the A allele of rs4251559
			with total serum IgE levels.			TLR2	rs13150331 rs4696480	of the IRAK4 gene are associated with high serum levels of IgE in the NP
							rs1898830 rs4696483	patients.
							rs3804100 rs5743704	
							rs5743708 rs7656411	
							rs1339 rs17030340	
							rs2289318 rs7695605	
						TLR3	rs956239 rs4861699	-
							rs5743305 rs7657186	
							rs6552950 rs3775296	
							rs35140061 rs7668666	
							rs3775292 rs35311343	
							rs5743317 rs3775291	
							rs5743318 rs10025405	
							rs4862633 rs4608848	
							rs6857595 rs1519309	
						TLR4	rs10983754 rs10759930	
							rs10759932 rs2149356	
							rs4986790 rs4986791	
							rs11536889 rs11536898	
							rs1554973 rs7860896	
							rs/037225 rs2183016	
						TLR6	rs5/43810 rs5743808	
							rs5/43/94 rs5/43/88	
							rsb833914	

		[[TI R9	rs352162 rs352140	
						TENS	13552102 13552140	
							rs5743836 rs187084	
							rs352143 rs11717574	
						TLR10	rs4513579 rs11466657	
							rs11096955 rs11466652	
							rs10856839 rs7653908	
						CD14	rc7701577 TC	
						0014	13//215//_10	
							rs4914_CG	
							rs2569190_GA	
							rs2569193_GA	
							rs2563310_GA	
						MD2/LY9	rs1905045_TC	
						6	rs16938755_TC	
							rs11786591_CT	
							rs6472812_GA	
							rs10504554_TC	
							rs17226566_TC	
							rs12544736_TG	
							rs16938766_GC	
						MYD88	rs2239621 rs4988453	
							rs7744 rs6767684	

							150790045	
						IRAK4	rs11182250 rs1461567	
							rs4251580 rs4251520	
							rs4251559 rs17121283	
							rs6582484 rs4251459	
							620-1delAC rs4251487	
							821delT rs4251583	
							T877C	
							134231313	
							A1188+520G	
							G1189-1T	
							rs4251545	
						TRAF6	rs3740961 rs5030437	
							rs5030416 rs5030411	
							rs331455	
Tournas et	 GWAS	Canada	To verify an association	196 HC	Blood	P73	rs3765731	The A allele of rs3765731 was
al. 2010	G 11/13	Cundud	between p73 and CRS.	454 000 100	5.000	,,,,,		differentially expressed in NP when
[153]				154 CRSwNP			123/02092	compared to no (p=0.037).
				52 CRSsNP				The A allele has a protective effect: AA+AG vs GG OR 0.5391, p=0.0036.
Wang et al.	CG	USA	To determine whether	123 HC	Blood	CFTR	ΔF508	Only 11 patients had one of the selected
2000 [68]			mutations in the CFTR	147 CBS			G542X	mutations in the CFTR gene.
			responsible for CF,	177 013			5572A	
			preuispose to CKS.				N1303K	
Wang et al. 2008 [130]	CG	Taiwan	To investigate the role of MMP2 tagging SNPSs	136 HC	Blood	MMP2	rs2438656 rs857403	rs857403 T allele was associated with increased risk (OR 2.07 p=0.03) but it
-			and promoter functional polymorphism in the	136 CRSwNP			rs1030868 rs1477017	could not be replicated with additional controls.

	1	1	I.		I.	r		4050605 0000674	
				development of NP.				rs1053605 rs9302671	
								rs2241145 rs2241146	
								rs243849 rs12599775	
								rs243847 rs243844	
								rs243840 rs2287076	
								rs11639960 rs243832	
								rs7201	
Wang et al.		CG	Taiwan	To investigate the role	730 HC	Blood	MMP9	rs3918242	The T allele of promoter SNP rs3918242
2010 [131]				and promoter functional	203 CRSwNP			rs2664538	dominant (nominal $p = 0.023$, empirical $p = 0.022$, $OR = 1.62$) and additive models
				development of NP.				rs3787268	(nominal p = 0.012, empirical p = 0.011, OP = 1.60) The A allele of re2274756 bac
								rs2274756	a nominal p value of 0.034 under the
									additive model, the most significant
									hapiotype was IGGA p=0.0045
Wang et al. 2013 [132]		CG	Taiwan	To investigate the relative expression of	31 HC	Blood	MMP2	rs243865	Genetic polymorphisms of MMP-2 and MMP-9 functional promoters were not
				MMPs in the non- recurrent and recurrent	30 CRSwNP		MMP9	rs3918242	associated with the recurrence of NP.
				NP as compared to control individuals.					
Yazdani et		CG	Iran	To investigate the	87 HC	Blood	CD14	rs946564423	Significant association of the C allele in
al. 2012 [133]				association between the polymorphism C-159T in	107 CRSwNP				NP patients (or 1.88, p=0.04)
()				CD14 gene and NP.					
Yea et al.		CG	Korea	To investigate the	70 HC	Blood	IL4	-590C/T	The presence of T allele was associated
2006 [45]				IL-4 promoter	106 CRS				p=0.028)
				polymorphism and NP.	61 CRSwNP				
Zhai et al.		CG	China	To explore a potential	81 HC	Blood	HLA-DR	*04	Frequency of allele was significantly
2007 [26]				association between NP and polymorphisms at	30 NP			*07	higher in patients for DR*09 and -*16 and DQ-*08 and -*09. DQ*07 frequency was
				loci for HLA-DR and HLA-DQ.				*08	lower in patients.
								*00	
								~09	
								*10	
								*11	
								*12	
								*13	
								*14	
								*15	
								*16	
							HLA-DQ	*02	
								*04	
								*05	
								*06	
								*07	
								*00	
								*08	
								*09	
Zhang et al. 2008 [134]		CG	China	To examine whether there is an association	180 HC	Blood	CC10 (SCGB1A1	+38A>G	No association
				between Clara cell 10kDa protein	90 CRSwNP)		
				(CC10)+38A>G, plasma CC10 levels and CRS.	130 CRSsNP				
Zhang -t -l			Canad-	To determine whether	197.00	Dia - d	NOST	**1004255	
2nang et al. 2011 [135]		CG.	Canada	polymorphisms in gene	181 HC	RIOOD	NUS1	151004350	p=0.0023, OR 0.62;
				regulating NO synthesis	154 CRSwNP			rs1483757	rs9658281, p =0.0129, O 0.66) remained

				are associated with CRS.	52 CRSsNP			rs545654	significant after correction for multiple
								rs9658281	testing. Homozygote allele C (p=0.0017; OR 0.28) in rs1483757 locus increased the risk.
							NOS1AP	rs10458392	rc12047527 in NOS1AB showed
								rs10919117	significant association (p<0.05) with CRS
								rs12022557	
								rs12047527	
								rs12061249	
								rs3923367	
								rs4657164	
								rs6676638	
								rs6677052	
								rs6677606	
								rs7416392	
								rs7546353	
								rs6681981	
								rc8179404	
Zhong et al			China	To configate and outpad	215.110	Blood	04053	*2072551	DedE22000 CND in DVDD increased the visk
2012a [40]		CG	China	genetic association	315 HC	BIOOD	PAR52	1528/3551	of CRSwNP (OR 2.76, p=3.2x10 ⁻⁶).
				Chinese population.	306 CRSWNP		IL22RA1	rs4292900	Selected SNPs in AOAH and IRAK4 were
					332 CRSSNP			rs4648936	0.60-0.79, p<0.05)
								rs16829225	
							TNFRSF1B	rs235214	
								rs496888	
								rs652625	
								rs7550488	
							TRIP12	rs1035833	
							IL1RL1	rs13431828	
								rs10204137	
		-					IL1A	rs17561	
								rs2856838	
								rs2048874	
								rs1800587	
							FAM79B	rs13059863	
							RYBP	rs4532099	
							TSLP	rs3806932	
							1521	rc2289276	
		-					10000	roje71E94	
							TNEAIDS	rc2757172	
							INFAIP3	rc5070038	
							141404	1372200	
							LAMBI	154/2/095	
							AOAH	rs4504543	
							MET	rs38850	
							RAC1	rs836479	
							CACNA2D 1	rs6972720	
							KIAA1456	rs11779957	
	1	1	1	1	1	1	1	1	1

		r		r			1	1
						MSRA	rs7001821	
						MUSK	rs10817091	
						PDGFD	rs12574463	
						NOS1	rs1483757	
						NAV3	rs1726427	
						IRAK4	rs4251431	
							rs6582484	
							rs1461567	
							rs3794262	
						SERPINA1	rs1243168	
							rs4900229	
						UBE3A	rs1557871	
						SLC13A3	rs393990	
						CACNA1I	rs3788568	
Zhang et al.	CG	China	To examine association	315 HC	Blood	EBI3	rs428253	Risk analysis showed rs428253 of EBI3
2013a [136]			between specific SNPs in/around the FOXP3	306 CRSwNP			rs6613	gene to play a protective role among both CRSsNP (GG/CC) and CRSwNP (CG/CC)
			and EBI3 genes and susceptibility to CRS	332 CRSsNP			rs353698	subjects. Haplotype analysis of the FOXP3 gene region further indicated that CRS
							rs2302164	risk was higher in individuals carrying the haplotype GG in rs2294018–rs2232365
						FOXP3	rs2294018	block, compared
							rs3060515	with wild-type AG haplotype
							rc2232365	
							rc3761548	
							rc4824747	
Zhang et al	 6	China	To explore associations	315 HC	Blood	TSID	rc1545169	SNPs rs252706 (AA genotype: P=0.012
2013b [137]	CG	China	between SNPs	206 CBSWND	ыооч	ISLF	rs754017	OR 0.552) and rs764917 (CA genotype: P=0.001_OR 0.182) displayed protective
			and development of CRS				13/0491/	roles among CRSwNP, but not CRScNP
				552 CR55INP			1512053750	subjects.
							1007 1000	
							rs12654933	
							rs10455025	
							rs11466741	
							rs13156086	
							rs6886755	
							rs252706	
							rs2416259	
Zielinska at al. 2012	CG	Poland	To investigate the association between LF	200 HC	Blood	LTF	rs1126478	Rs1126478 LF (OR 4.78; 95% CI 3.07– 7.24), the -33C/G OSF2 (OR 3.48; 95% CI
[138]			and OSF2 polymorphisms with the	195 CRSwNP		fgOSF2	rs3829365	2.19–5.52) and the rs3829365 OSF2 (OR 16.45; 95% Cl 6.71–40.30) genotypes
			risk of CRSwNP in Poland				-33C/G	were associated with an increased risk of CRSwNP.
1		1		1			1	1

Reference	Objective	Tissue	Epigenetic assay	Population	Significant findings
Callejas-Diaz et al. 2020 [84]	To identify which key mRNA and miRs are regulating in vitro mucociliary differentiation of human adult basal stem cells under pathological and healthy conditions.	NP, inferior turbinate mucosa (ITM; control)	miRNA	Spain	Transcriptome related to ciliogenesis and cilia function is significantly impaired during differentiation of CRSwNP epithelium due to an altered expression of microRNAs, particularly of those belonging to mir-34 and mi- 449 families
Cheong et al. 2011 [76]	To analyze the genome-wide DNA methylation levels in nasal polyps from patients with AIA.	NP, buffy coats	Genome-wide DNA methylation	China	 332 loci in 296 genes were hypermethylated in AIA vs ATA. These genes are involved in ectoderm development, hemostasis, and wound healing. 158 loci in 141 genes were hypomethylated in AIA vs ATA. Relevant pathways were lymphocyte proliferation, cell proliferation, leukocyte activation, and immune response.
Cho et al. 2012 [75]	To study the effect of trichostatin A (TSA), a histone deacetylase inhibitor, on TGFβ1-induced myofibroblast differentiation and ECM accumulation in NP fibroblasts.	NP, ITM	Histone acetylation control	Korea	The expression levels of HDAC2, α -SMA and TGF- β 1 were increased in NP compared to normal tissues. TSA induced hyperacetylation of histones, inhibiting them. HDAC inhibition is associated with myofibroblast differentiation and ECM accumulation in NP.
Cho et al. 2013 [75]	To investigate the inhibitory effect of TSA on myofibroblast differentiation and ECM production in nasal polyp organ cultures.	NP tissue cultures	Histone acetylation control	Korea	TSA inhibited HDAC and induced hyperacetylation of histones H4
Kidoguchi et al. 2018 [77]	To investigate the methylation levels at 3 CpG sites in the proximal PLAT promoter and their effects on gene expression in NP tissue.	NP, ITM	DNA methylation	Japan	Methylation of -618, -121, and -105 CpGs was significantly higher in NP. <i>PLAT</i> expression was lower (p>0.001). The methylation changes at -618 site showed a negative correlation with the gene expression changes between NP and ITM (r=65, p<0.01).
Kim et al. 2018 [78]	To elucidate whether DNA methylation of specific genes is involved in the development of NP.	NP, ITM	DNA methylation	Korea	The promotor regions of 10 and 30 genes were hypermethylated and hypomethylated, respectively, in NP samples compared with controls. The top four genes with altered hypomethylation in NP tissues were <i>KRT19, NR2F2, ADAMTS1</i> and <i>ZNF222</i> .
Kim et al. 2019 [79]	To investigate the expression and distribution of FZD5 and the role of eosinophil infiltration in CRSwNP pathogenesis.	NP, uncinated process tissue	Methylation profiling	Korea	397 and 387 genes were differentially hypermethylated and 399 and 208 genes were hypomethylated in the E- CRSwNP and NE-CRSwNP groups, respectively, compared to the control tissues. Most of the differentially methylated genes were associated with cancer pathways. FZD5 was significantly hypomethylated in the E- CRSwNP compared to the NE-CRSwNP group.
Li et al. 2019a [80]	To determine whether there was any association between abnormal DNA methylation of TSLP gene and CRS	NP, ethmoid mucosa (CRSsNP) patients	DNA methylation	China	There was an increase in methylation ratios of 4 CpGs (2, 22, 23, 24) of TSLP gene had increased in the CRSwNP patients compared to the CRSsNP and

	pathogenesis.	and ITM			control subjects, significantly related to disease status (p<0.02)
Li et al. 2019b [81]	To determine whether there was any association between abnormal DNA methylation of IL8 promoter and CRS pathogenesis.	NP, ethmoid mucosa (CRSsNP) patients and ITM	DNA methylation	China	Three CpGs (-116, -106, -31) were significantly hypomethylated in the CRSwNP group compared with CRSsNP and HC.
Liu et al. 2018 [85]	To study the role of miR124 in CRSwNP.	NP, ITM	miRNA	China	MiR124 expression was reduced in NP tissues, which negatively correlated with the expression of AHR. This may be critical to the development of inflammatory response in CRSwNP.
Liu et al. 2019 [83]	To characterize the transcriptome profiles of mRNAs and IncRNAs in patients with CRSwNP.	GEO datasets, blood samples	IncRNA	China	A total of 265 differentially expressed IncRNAs were obtained, including 56 upregulated and 209 downregulated genes.
Luo et al. 2017 [86]	To test whether miR-17-92 cluster is associated with suppressing IL-10 in peripheral DC.	Blood samples	miRNA	China	A negative correlation was found between expression of II-10 and miR- 19a in DC from NP patients. miR19-1 was upregulated while miR-17, -18a, - 19b, -20a and -92a showed no differences between NP and HC.
Ma et al. 2015 [88]	To investigate miRNAs expression profiles of peripheral blood DCs in CRS patients.	Blood samples	miRNA	China	There were 31 miRNAs changed in all CRS patients with respect to HC, and 49 miRNA that changed exclusively in CRSwNP. miR-210-3p, miR-125b-5p, and miR- 150-5p were upregulated in CRS, while miR-708-5p and miR-126-3p were downregulated.
Ma et al. 2018 [87]	To investigate the effects and mechanism of miR-150- 5p to promote the development of CRS via the DC-Th axis.	Blood samples	miRNA	China	miR-150-5p was upregulated in DCs from CRS patients compared with HC, and DCs Promote Naïve T Cells Proliferation. MiR-150-5p further regulated EGR2 and inhibited DCs, leading to an abnormal DC-Th axis.
Qing et al. 2019 [89]	To investigate the mechanisms between the miR-142-3p and TNF-a activation in vitro and in vivo	NP, ITM	miRNA	China	miR-142-3p may participate in the regulation of the body's inflammatory response through the LPS-TLR-TNF-a signaling pathway in CRSwNP.
Seiberling et al. 2012 [95]	To determine the presence of 5-bromo-cytosine, 5- chloro-cytosine and methylated cytosine in CRSwNP.	NP, posterior ethmoid tissue (HC)	DNA modification	USA	The levels of 5-Bromocytosine were significantly higher in polyps (p=0.007). Aberrant methylation patterns in polyp eosinophils may help explain the pathogenesis of CRSwNP.
Tian et al. 2012 [96]	To explore the profiling of tandem alternative polyA (APA) sites in NP.	NP, uncinated process mucosa	Genome-wide polyadenylation site sequencing	China	There was a switching of 3'UTR lengths in NP compared with nasal uncinate process mucosa from the same patient. 105 genes were switched to distal polyA sites in the nasal polyps and 90 genes were switched to proximal poly(A) sites. Besides, 213 genes were upregulated in NP while 414 genes were downregulated.
Xuan et al. 2019 [90]	To evaluate miRNAs profiles and relevant biological pathways in CRSwNP and	Nasal mucosa	miRNA array	China	24 miRNAs showed differential expression. 5 miRNAs (miR-210-5p, miR-3178 miR-585-30 miR-3146 and

	control subjects.				miR-320e) were significantly upregulated (p < 0.05, fold change >2), and 19 miRNAs, including miR-32-3p, miR-1299, miR-3196, miR-3924, miR- 548e-3p, miR-3184-5p, miR-375, miR- 23a-5p, miR-377-5p, miR-574-5p, miR- 3149, miR-500a-5p, miR-125b-2-3p, miR-1914-5p, miR-532-3p, miR-612, miR-1298-5p, miR-1226-3p, and miR- 668-3p, were significantly downregulated in CRSwNP tissue (p < 0.05, fold change <0.5).
Yan et al. 2020 [91]	To examine human neutrophil elastase-induced MUC5AC overexpression in CRS via miR-146a.	NP, uncinated process mucosa	miRNA	China	EGFR is a target of miR-146a. This miRNA is downregulated in NP reducing the inhibition of EGFR, and therefore MUC5AC expression levels were increased.
Yu et al. 2018 [92]	To evaluate the roles of TGFβ1 and miR-663 in the pathogenesis of NP in children.	Nasal mucosa, peripheral blood eosinophils (PBE)	miRNA	China	The expression of miR-633 was significantly reduced in polyps and PBE from CRS patients, while <i>TGFB1</i> mRNA was significantly increased. miR-633 binds to the 3'UTR of <i>TGFB1</i> and regulated its expression.
Zhang et al. 2012b [94]	To determine the pattern of expression and biological role of miRNAs in CRS.	NP, ethmoidal mucosa, inferior turbinate tissue	miRNA	China	miR-125b was upregulated in CRSwNP when compared to CRSsNP. This may enhance type I IFN expression through suppressing 4E-BP1 protein expression in airway epithelial cells.
Zhang et al. 2012c [97]	To investigate the expression of miRNA machinery components in CRS.	NP, ethmoid sinus mucosa	mRNA expression	China	PACT mRNA expression was found to be upregulated in CRSwNP as compared with controls. The rest of the miRNA machinery components including Drosha, Dicer, TRBP, FXR1 and E1F2C2, showed no differences between patients and controls.
Zheng et al. 2015 [82]	To identify whether DNA methylation pays a role in the pathogenesis of NP.	NP, ITM	DNA methylation	China	198 genes had a differential methylated signal in their promoter region when comparing NP samples with ITM samples. The four most changed genes were <i>COL18A1</i> , <i>EP300</i> , <i>GNAS</i> and <i>SMURF1</i> .
Zhou et al. 2020 [93]	To explore the pathogenesis of CRSwNP from the perspective of genes.	CRSwNP datasets. NP, nasal mucosa (HC)	Functional enrichment analysis, including non- coding RNAs	China	Two clusters of genes, IncRNAs and miRNAs were found to be related to CRSwNP. Main miRNA involves were: miR-130a, miR-27a-3p, miR-193-3p, miR-29a-3p, miR-18b-5p, miR-138-5p, and miR-25- 3p.

Functional Category	Enrichment FDR	Genes in list	Total genes	Genes
Cytokine-mediated signaling pathway	2.63e-16	29	950	IL1B IL1RN IL22RA1 CCL11 IRAK4 TSLP EBI3 IL1RL1 FCER1G IL1A PPARG TNF NOS2 ALOX5 MMP2 MMP9 IL10 IL33 ALOX15 CIITA HLA-DRB3 HLA-DRB1 HLA- DQA1 HLA-DRB5 HLA-DRA HLA-C HLA-B HLA-A HLA-DRB4
Defense response	1.10e-15	38	2062	NOS2 IL33 FCER1G PTGDR CD14 CCL11 CIITA LTF IL1B IL10 TNF HLA-A ALOX5 FOXP3 IL1A IL1RL1 PPARG ALOX5AP AOAH IL1RN IL22RA1 MS4A2 ADORA1 CYSLTR1 SERPINA1 IRAK4 AGER TSLP HLA- DRB1 MMP9 ALOX15 HLA- DRB3 HLA-DQA1 HLA-DRB5 HLA- DRA HLA-C HLA-B HLA-DRB4
Inflammatory response	1.10e-15	27	856	IL33 PTGDR CD14 CCL11 CIITA NOS2 IL1B IL10 TNF ALOX5 FOXP3 IL1A IL1RL1 PPARG ALOX5AP AOAH IL1RN MS4A2 FCER1G ADORA1 CYSLTR1 SERPINA1 AGER TSLP HLA-DRB1 MMP9 ALOX15
Response to stress	7.689e-15	52	4507	NOS2 MMP2 CAT IL1B HSPA2 IL1RN IL33 TRIP12 FANCC FCER1G PTGDR CD14 CCL11 MSRA CIITA CFTR LTF DCBLD2 TP73 NOS1 MMP9 IL1A IL10 TNF HLA-A IFRD1 ALOX5 FOXP3 IL1RL1 PPARG ALOX5AP AOAH IL22RA1 MS4A2 ALOX15 ADORA1 ADRB2 CYSLTR1 SERPINA1 IRAK4 MT-CO2 AGER MET TSLP HLA- DRB1 HLA-DRB3 HLA-DRA HLA-C HLA-B HLA-DRB4

Response to cytokine	1.04e-13	30	1372	IL1B IL1RN
				IL22RA1 CCL11
				IRAK4 TSLP CIITA
				NOS2 EBI3
				IL1RL1 FCER1G
				MMDg II 10 II 33
				DRB1 HLA-DOA1
				HLA-DRB5 HLA-
				DRA HLA-C HLA-B
				HLA-A HLA-DRB4
Immune system	1.124e-13	45	3539	RUNX2 IL1B
process				IL1RN CD8A
				FCER1G CD14
				CCL11 ACE LTF
				HIA-R HIA-A
				NOS2 MMP9 FBI3
				TAPBP IL1A
				IL1RL1 MS4A2
				FANCC ALOX15
				ADORA1
				CYSLTR1 LTA
				HLA-DQB1 HLA-
				DRB3 HLA-DQA1
				IRAK4 HLA-DRB5
				TD73 CAT
				FCFR1A
				SERPINA1
Cellular response to	1.121e-13	29	1278	IL1B IL1RN
cytokine stimulus				IL22RA1 CCL11
				IRAK4 TSLP
				NOS2 EBI3
				IL1RL1 FCER1G
				MMPG II 10 II 33
				DRB1 HLA-DQA1
				HLA-DRB5 HLA-
				DRA HLA-C HLA-B
				HLA-A HLA-DRB4
Immune response	1.41e-13	39	2602	CD8A FCER1G
				CD14 CCL11 IL1B
				IL1RN IL33 MS4A2
				ALOX15 CYSLTR1
				LTA HLA-DQB1
				HLA-DRB3 HLA-
				DRB1 HLA-DQA1
				IRAK4 HLA-DRB5
				MMP9 FRIS
				TAPBP CAT
				FCER1A
				SERPINA1
Cell surface receptor	2.29e-13	43	3287	MUSK MET IL1B
signaling pathway				IL1RN IL22RA1
		1	1	ANAA4 IRAN4 LI F

				TSLP TNF FOXP3
				FBI3 II 1A II 1RI 1
				RUNX2 33
				MS4A2 ADORA1
				ADRB2 CVSI TR1
				MMP9 PPARG
				AGER HLA-C
				HLA-B HLA-A
				HLA-DRB4
Cellular response to	4.85e-13	44	3536	IL1B HSPA2
chemical stimulus				PPARG IL1RN
				IL22RA1 FANCC
				FCER1G CD14
				CCL11 MSRA
				IRAK4 CFTR LTF
				MMP9 ALOX5AP
				TSLP ALOX15
				AGER TNF NOS2
				MMP2 NOS1 EBI3
				II 1RI 1 CAT
				RUNX2 10
				PTGDR ADRB2
				CIITA I TC4S MET
	5 00 40			
Cellular response to	5.90e-12	39	2938	IL1B HSPA2
organic substance				PPARG IL1RN
				IL22RA1 CD14
				CCL11 IRAK4
				CFTR LTF TSLP
				AGER TNF NOS2
				MMP2 NOS1 EBI3
				IL1RL1 CAT
				RUNX2 IL10
				FCER1G PTGDR
				ADRB2 CIITA IL1A
				ALOX15 HLA-
				DRB1 ALOX5
				MMP9 IL33 HLA-
				DRB3 HLA-DQA1
				HLA-DRB5 HLA-
				DRA HLA-C HLA-B
				HLA-A HLA-DRB4
Response to organic	1.593e-11	42	3547	NOS2 NOS1 IL1B
substance				HSPA2 PPARG
				IL1RN IL22RA1
				CD14 CCL11
				IRAK4 CFTR I TF
				IL10 TSLP HI CS
				CIITA AGER TNE
				TBXAS1 TP73
				MMP2 MMP0 FRI3
				RUNX2 FCFR1G
				HLA-C HLA-B
				HLA-A HLA-DRB4
Regulation of immune	2.414e-10	30	1909	FCER1G CD14
	1		1	

				10 33 AGER
				TNF HLA-B HLA-A
				IL1RL1 MS4A2
				ALOX15 ADORA1
				LTF PPARG TSLP
				HLA-DRB1 TP73
				EBI3 CD8A
				FCER1A HLA-
				DRB3 HLA-DQA1
				IRAK4 HLA-DRB5
				HLA-DRA HLA-C
				HLA-DRB4
Regulation of immune	5.63e-10	25	1325	FCER1G CD14
response				IL1B FOXP3 IL10
				AGER TNF HLA-B
				HLA-A IL1RL1
				IL33 MS4A2
				ALOX15 LTF
				PPARG CD8A
				FCER1A HLA-
				DRB3 HLA-DRB1
				HLA-DQA1 IRAK4
				HLA-DRB5 HLA-
				DRA HLA-C HLA-
				DRB4
Regulation of response	1.05e-09	46	4820	IL1B IL1RN IL33
to stimulus				FCER1G CD14
				CCL11 IRAK4
				NOS2 LTF FOXP3
				MET IL10 TSLP
				ADRB2 AGER
				TNF HLA-B HLA-A
				CFTR TP73 NOS1
				EBI3 IL1A IL1RL1
				CAT RUNX2
				PPARG ALOX5AP
				AOAH MS4A2
				ALOX15 ADORA1
				LTA RGS7BP
				MMP9 TRIP12
				HLA-DRB1
				NOS1AP CD8A
				FCER1A HLA-
				DRB3 HLA-DQA1
				HLA-DRB5 HLA-
				DRA HLA-C HLA-
				DRB4
Cvtokine secretion	3.78e-09	13	285	CD14 NOS2
-,		-		FOXP3 IL1A IL10
				IL33 TNF IL1RL1
				IL1B AGER TSLP
				ANXA4 HLA-DRB1
Cellular response to	4.23e-09	13	289	CCL11 NOS2
interferon-gamma				CIITA PPARG
				HLA-DRB3 HLA-
				DRB1 HLA-DQA1
				HLA-DRB5 HLA-
				DRA HLA-C HLA-B
				HLA-A HLA-DRB4
Interferon-gamma-	4.76e-09	11	178	PPARG CIITA
mediated signaling				HLA-DRB3 HLA-
pathway				DRB1 HLA-DQA1
				HLA-DRB5 HLA-
				DRA HLA-C HLA-B
				HLA-A HLA-DRB4
Regulation of	6.81e-09	15	448	IL33 NOS2
inflammatory response	-			FOXP3 IL1RL1
				IL1B PPARG
				ALOX5AP AOAH
				IL10 FCER1G
				ADORA1 AGER
				TSLP TNF MMP9
Antigen processing and	9.27e-09	14	384	FCER1G HLA-
presentation				DRB1 HLA-DRA
-				HLA-B TAPBP

				HLA-DQB1 HLA-
				DRB3 HLA-DQA1
				HLA-DRB5 HLA-C
				HLA-A HLA-DRB4
Deenenge to interferen	0.070.00	10	210	
Response to interferon-	9.276-09	15	312	
gamma				
				DRA HI A-C HI A-B
				HI A-A HI A-DRB4
Regulation of cytokine	1.269e-08	12	257	CD14 FOXP3
secretion				IL1A IL10 IL33
				TNF IL1RL1 IL1B
				AGER TSLP
				ANXA4 HLA-DRB1
Positive regulation of	1.51e-08	32	2621	IL1B IL1RN IL33
response to stimulus				FCER1G CD14
				CCL11 IRAK4 LTF
				IL10 TSLP ADRB2
				TNF HLA-B CFTR
				FOXP3 TP73
				NOS1 IL1RL1 CAT
				ALOX5AP
				ADORA1 AGER
				ALOX15 HLA-
Cytokine production	4 970-08	10	925	
Cytokine production	4.376-00	19	920	NOS2 TE EOXP3
				TSI P AGER TNF
				IL1RL1 FCER1G
				IRAK4 NAV3
				ANXA4 HLA-DRB1
				EBI3
Interleukin-6 production	4.97e-08	10	172	IL1B IL1RN NOS2
				IL10 TNF FOXP3
				IL33 FCER1G
				AGER TSLP
Secretion	7.35e-08	26	1861	CACNA1I CD14
				NOS2 FOXP3 IL1A
				IL1B IL10 IL33
				ACE TNF CFTR
				IL1RL1 IL1RN
				FCER1G ADORA1
				MI-CO2 AGER
				ISLP ANXA4 HLA-
				DRB1 LTF ALOX5
Positive regulation of	7 350 09	20	1069	
transport	7.350-08	20	1069	
transport				
				PPARG ADORA1
				AGER ADRB2
				TSLP NOS1 HI A-
				DRB1 NOS1AP
				TP73
Regulation of cvtokine	8.36e-08	18	852	IL1B IL1RN CD14
production				NOS2 LTF FOXP3
				IL1A IL10 IL33
				TSLP AGER TNF
				IL1RL1 FCER1G
				NAV3 ANXA4
				HLA-DRB1 EBI3
Regulation of defense	8.789e-08	19	968	IL33 CD14 NOS2
response				HLA-A FOXP3

Supplementary Table 3.

				IL1RL1 IL1B PPARG ALOX5AP AOAH IL10 FCER1G ADORA1 AGER LTF TSLP TNF MMP9 IRAK4
Regulation of peptide secretion	9.132e-08	15	559	CD14 FOXP3 IL1A IL1B IL10 IL33 TNF CFTR IL1RL1 ADORA1 AGER NOS2 TSLP ANXA4 HLA-DRB1