Asthma is a complex and multifactorial respiratory disease with a high prevalence in the pediatric population. Variation in treatment response to asthma therapies has been described among patients, and difficult-to-treat asthma carries both high healthcare and socioeconomic burden to the patients and society. Omic studies can be used to discover the molecular mechanisms underlying asthma susceptibility and treatment response, contributing to a better knowledge and definition of asthma pathogenesis and therefore, to the development of precision medicine. This entry aims to summarize the recent findings of omic studies of treatment response in childhood asthma. Between 2018-2019 a total of 13 omic studies has been performed involving genomics, epigenomics, transcriptomics, metabolomics, and the microbiome. These have been focused on the response to three common asthma medications: short-acting beta agonists, inhaled corticosteroids, and leukotriene receptor antagonists. Novel associations of different biomarkers with asthma treatment response have been described. However, stronger evidence and more consistent results are required to implement these molecular biomarkers into clinical practice by establishing the most appropriate therapy for each patient.

**Introduction**

Asthma, the most prevalent chronic respiratory disease in children, is a heterogeneous and complex disease characterized by airway inflammation, reversible airflow obstruction, and bronchial hyperreactivity. The most characteristic asthma symptoms are wheeze, cough, chest tightness, and shortness of breath. These symptoms may be prevented, relieved, and controlled by common asthma medications, such as short-acting beta agonists (SABAs), inhaled corticosteroids (ICSs), or leukotriene receptor antagonists (LTRAs). However, some patients do not achieve adequate control of the disease even with the most intense therapies, such as ICSs at high-doses or oral corticosteroids (OCSs). These patients are more prone to develop asthma exacerbations, worse episodes of the disease which carry a burden to the patients, their families, and the community. Different phenotypes and endotypes of asthma have been described, each one with both shared and differentiated features including the response to pharmacological therapies. Nonetheless, phenotype-guided treatments are scarce and a clear relationship between asthma phenotypes and treatment response has not been established. In this context omic sciences, which consist of high-throughput analyses at different biological layers, would aid to disentangle the molecular mechanisms underlying asthma and treatment response to move toward precision medicine. This entry aims to summarize the findings of omic studies of asthma treatment response conducted between 2018-2019 in pediatric populations (Figure 1).
**Pharmacogenomics**

Pharmacogenomics investigates the association of genetic markers across the genome with treatment response. This approach usually focuses on single nucleotide polymorphisms (SNPs), which are changes in the DNA sequence affecting one nucleotide shared by >1% of the population. This type of marker is usually co-inherited with other genetic variants due to the linkage disequilibrium (LD) patterns allowing the indirect study of millions of genetic variants that are inferred from hundreds of thousands of SNPs assessed by genotyping arrays. Thus, common genome-wide genetic variation can be studied without any prior hypothesis by genome-wide association studies (GWAS) \[6\]. Recently, whole-genome sequencing (WGS) emerged as a high-resolution method to study both common and rare genetic variation. Although WGS detects genetic variation not tackled by genome-wide genotyping arrays, its high cost has still limited its application to pharmacogenomic studies \[7\].

Recent pharmacogenomic studies of childhood asthma have not been limited to European-descent populations as in the past \[8\]–\[9\], but they have mostly analyzed two minority populations from the United States with a high prevalence of asthma and low treatment response \[10\]: African Americans and Hispanics/Latinos. Specifically, several publications within the period reviewed have been focused on two studies: Study of African Americans, Asthma, Genes and Environments (SAGE) and the Genes-Environment and Admixture in Latino Americans (GALA II) (Table 1). In these studies, genomic analyses of SABA treatment response were carried out by means of GWAS using genotyping arrays and WGS data \[11\]–\[12\]. Additionally, given that African Americans and Hispanics/Latinos are admixed populations, admixture mapping can be applied in order to identify genomic regions in which local ancestry is associated with pharmacogenetic traits \[12\]. This complementary approach is possible because of the differences in allele frequency of the SNPs depending on their ancestral background \[10\]. These studies have been focused on bronchodilator drug response (BDR), a measurement of the change in lung function after SABA administration \[11\]–\[12\], and also on the presence/absence of severe asthma exacerbations despite ICS treatment as a proxy of ICS response \[13\].

**Table 1.** Summary of the genomic regions identified by pharmacogenomic studies in childhood asthma performed during 2018-2019.

| rsID/Chr. Band | Chr:Positiona | Gene/Nearest Gene | Effect Allele | Effect | p-value | Reference |
|----------------|---------------|-------------------|---------------|--------|---------|-----------|
| rs73650726     | 9:85152666    | 9q21              | A             | $\beta = -3.8$ | $7.69 \times 10^{-9}$ |           |
| rs7903366      | 10:53689774   | PRKG1             | T             | $\beta = 1.23$  | $3.94 \times 10^{-8}$ |           |
Spear et al. performed a GWAS of BDR in African Americans with asthma from SAGE (n=949) analyzing imputed data from genotyping arrays. A SNP from 9q21 (rs73650726) was genome-wide significantly associated with BDR exclusively in African-admixed populations. Moreover, a trans-ethnic meta-GWAS across African American and Hispanic/Latino children and youth (n=2779) identified genome-wide associations of three SNPs from the protein kinase cGMP-dependent 1 (PRKG1) gene with SABA response. Interestingly, PRKG1 is a biologically plausible gene for SABA response given that it encodes a protein involved in the nitric oxide/cGMP signaling pathway, acts as an important regulator of airway inflammation in response to SABA, and participates in smooth muscle relaxation.

Furthermore, PRKG1 has been associated with lung function measurements and asthma risk. However, these results need to be replicated in independent studies.

A WGS study was conducted in the same populations selecting patients with asthma with extreme BDR values among African Americans, Puerto Ricans, and Mexicans (n = 1441). This revealed two SNPs located near the dynein axonemal heavy chain 5 (DNAH5) gene as significantly associated with BDR in a trans-ethnic meta-analysis. DNAH5 is ATPase protein that participates in a complex associated with the microtubules. Variants of this gene have been associated with allergic sensitization, lung function, and immunoglobulin E (IgE) serum levels. The WGS study by Mak et al. also revealed population-specific associations of both common and rare variants (1p13.2 and 11p14.1 in Mexicans and 19p13.2 in African Americans) (Table 1).

Finally, an additional study on the GALA II and SAGE populations evaluated the ICS response in a subset of Hispanic/Latino and African American asthma patients treated with ICSs. ICS response was defined as the presence/absence of asthma exacerbations in the past 12 months despite ICS treatment (Table 1). Asthma exacerbations were defined by any of the following events originated from asthma symptoms: emergency room (ER) visits, hospitalizations, or OCS use. The
association of a SNP from the intergenic region of the genes encoding for the apolipoprotein B mRNA editing enzyme catalytic subunits 3B and 3C (APOBEC3B-APOBEC3C) was revealed, a result that was replicated in European children with asthma (n = 1697). However, this SNP did not reach the genome-wide significance threshold in a meta-analysis across all populations. Despite this, a consistent association of the SNP from the APOBEC3B-APOBEC3C intergenic regions was found with the change in the forced expiratory volume in the first second (FEV1) after ICS treatment during 6 weeks in a subset of patients with such data available (n = 166), as an alternative measurement of ICS response. APOBEC3B and APOBEC3C are subunits of a cytidine deaminase that carries out RNA editing and participates in the immune response to viruses. In the same study [13], an attempt of replication of genomic regions previously associated with ICS response was performed. The association of the region that included the L3MBTL4 and ARHGAP28 was validated, which has also been associated with post-bronchodilator lung function [18].

**Epigenomics**

Epigenetics is the study of the mechanisms involved in gene expression regulation without modifying the DNA sequence. These include DNA methylation (DNAm), microRNA (miRNA) regulation, and histone modifications. Nowadays, whole-genome epigenetic changes can be analyzed through high-throughput techniques. DNAm is the most studied epigenetic modification, involving the methylation of a cytosine base followed by a guanine (CpG site) [21][22][23].

Two epigenome-wide association studies (EWAS) were performed during the period reviewed evaluating the effects of ICS response on DNA methylation patterns in peripheral blood cells (PBCs) [24][25]. In the first study [24], non-Hispanic whites from the Childhood Asthma Management Program (CAMP, n = 154), Europeans from the Children, Allergy, Milieu, Stockholm, Epidemiology (BAMSE, n = 72), and Hispanic/Latinos from the Genetic Epidemiology of Asthma in Costa Rica Study (GACRS, n = 168) were analyzed. Treatment response was measured as two outcomes related to asthma exacerbations in the past 12 months while the patients were treated with ICSs. Hypomethylation of a CpG site near interleukin 12B (IL12B) was protective for severe exacerbations in a meta-analysis across non-Hispanic whites and Hispanic/Latinos, and this hypomethylation was correlated with lower IL12B expression in blood cells in one of the studies. IL12B encodes a subunit of two cytokines (IL-12 and IL-23) involved in immune response and airway hyperresponsiveness, and its expression level has been associated with corticosteroids response in bronchial biopsies from asthma patients [26][27][28]. Moreover, in the same study, Wang et al. found 13 CpG sites that were significantly associated with the absence of OCS use in a meta-analysis across non-Hispanic whites (n = 322). An interaction analysis identified that hypermethylation of a CpG near the genes encoding for the cortistatin (CORT) and centromere protein S (CENPS) was associated with the absence of OCSs use, and also with higher CORT expression in CAMP [26][27][28]. Interestingly, cortistatin is a peptid involved in the anti-inflammatory process regulated by the hypothalamic-pituitary-adrenal axis and regulates endogenous corticosteroid levels [22][30].

**Table 2.** Summary of the main findings of epigenomic studies focused on DNAm in childhood asthma performed during 2018-2019.

| CpG       | Chromosome:Positiona | Gene/Nearest Gene | β      | p-value | Reference |
|-----------|----------------------|-------------------|--------|---------|-----------|
| cg00066816 | 5:158758353          | *IL12B*           | −3.101 | 0.002   |           |
| cg00557354 | 13:111767899         | *ARHGEF7*         | −3.490 | 0.001   |           |
| cg04256470 | 1:10510465           | *CORT CENPS*      | 3.620  | <0.001  |           |
| cg09495977 | 4:8271507            | *HTRA3*           | −2.420 | 0.017   |           |
| cg12333095 | 12:110437035         | *ANKRD13A*        | −3.485 | 0.001   |           |
The second EWAS of ICS response was performed in non-Hispanic white children from CAMP (n = 152) analyzing the change in FEV₁ after eight weeks of ICS treatment[25]. Relative hypermethylation of a CpG from the bolA family member 2 (BOLA2) gene was associated with lung function improvement and with increased gene expression of BOLA2. BOLA2 is involved in the maturation process of iron-sulfur containing proteins[31]. Gene expression levels of BOLA2 in airway cells differ by asthma status[29], and intronic variants of this gene are associated with eosinophil levels and lung function measurements[17][32][33]. Additionally, the study by Wang et al. validated the hypermethylation of the orthodenticle homeobox 2 (OTX2) gene in PBCs, which had been previously associated with good response to OCS in nasal cells[34].

Regarding miRNA profiling without a prior hypothesis, one study has been performed between 2018-2019. Kho et al. evaluated the association of 754 circulating miRNAs in serum with ICS response in non-Hispanic white children with asthma from CAMP (n = 153)[35]. A total of 12 miRNAs were associated with the risk of exacerbations, defined as steroid bursts in the past year because of asthma ICS treatment (Table 3). The strongest association was described for miR-206, which had higher serum expression levels in non-exacerbators compared to exacerbators. In addition, a predictive model for asthma exacerbations based on three miRNAs (miR-206, miR-146b-5p, and miR-720) and clinical variables had an improved prediction area under the receiver characteristics curve (AUC) than a model exclusively based on clinical parameters (0.81 vs 0.67). Strikingly, these three miRNAs have been related to asthma in cellular and animal models[33], and miR-146b-5b and miR-206 had also been associated with baseline FEV₁/FVC[36]. Moreover, four biological pathways regulated by the three miRNAs were revealed by Kho et al., including two previously associated with asthma[35].

Table 3. Summary of findings of the only epigenomic study focused on the association of circulating miRNA with inhaled corticosteroid (ICS) response in childhood asthma during 2018-2019[35].

| miRNA   | Odds Ratio | p-value |
|---------|------------|---------|
| miR-206 | 0.60       | 0.004   |
| miR-146b-5p | 0.66   | 0.007   |
| miR-222-3p | 0.70   | 0.02    |
Transcriptomics

Transcriptomics is the study of all RNA transcripts expressed in a cell or individual by high-throughput methods such as RNA sequencing (RNA-seq) or microarrays. During 2018-2019, most transcriptomic studies of treatment response in pediatric asthma focused on ICS response. A microarray analysis of peripheral blood mononuclear cells (PBMCs) from Taiwanese children with asthma (n = 133) showed that patients with poor asthma control have a specific transcriptomic pattern associated with glucocorticoid signaling and immune response [37]. Moreover, the application of system biology approaches in the analysis of transcriptomic data of ICS response in non-Hispanic white children from CAMP has been performed by two studies. In the first study, Qui et al. [38] analyzed gene expression networks in immortalized B-cell lines from children treated with ICSs during two months and classified as good (n = 47) or poor (n = 48) ICS responders based on changes in lung function after ICS treatment. While good responders had an enrichment in immune response and proapoptotic corticosteroid-induced pathways, poor responders showed enrichment in antiapoptotic pathways. Additionally, two transcription factors (TFs), the nuclear factor kappa B subunit 1 (NFKB1) and the Jun proto-oncogene, AP-1 transcription factor subunit (JUN), had differential regulation between both groups. Interestingly, dysregulation of NFKB1 is associated with inflammatory diseases and inadequate immune cell development, and JUN has been related to macrophage activation [39].

McGeachie et al. [40] conducted a multi-omic analysis in non-Hispanic white children with asthma from CAMP treated with budesonide (n = 104). Treatment response was evaluated by a composite phenotype of ICS response, which was integrated with genome-wide genotyping data and the response of immortalized lymphoblastoid cells to dexamethasone into a network. A total of seven genes were associated with steroid response and four of them were selected for in vitro functional analyses in lung epithelial cells. A knockdown of the family with sequence similarity 129 member A (FAM129A) gene showed a reduced response to dexamethasone. FAM129A encodes for a protein involved in an apoptosis pathway [41].

Katayama et al. [42] performed a transcriptomic study of LTRA response in children aged 6–48 months recruited during an acute wheezing episode and followed-up to seven years (n = 107). A weighted gene co-expression network analysis (WGCNA) [43] was performed in leukocytes to identify modules of genes involved in LTRA response. A module of 145 co-regulated genes involved in interferon signaling pathways, inflammation, and antiviral response was correlated with acute wheezing. Moreover, this module was associated with lung function and the number of asthma exacerbations, and had a good predictive value for future LTRA medication (AUC = 0.81). The tripartite motif containing 22 (TRIM22) gene showed the strongest association with LTRA treatment. Interestingly, TRIM22 is involved
Metabolomics provides the profile of the whole metabolite composition (metabolome) in biological samples by means of high-throughput analytical techniques, such as mass spectrophotometry, nuclear magnetic resonance, or spectroscopic methods. Interestingly, metabolomics can be performed in non-invasive samples such as exhaled air (breathomics) and this recent omic has been applied to asthma with promising results \([45][46][47]\).

Kelly et al. examined the interaction of age and 501 serum metabolites on BDR after albuterol administration in blood samples from children with asthma from CAMP at three-time points every four years, with mean ages of 8.8 \((n = 560)\), 12.8 \((n = 563)\), and 16.8 \((n = 295)\), respectively \([48]\). A total of 39 metabolites, mainly lipids, showed a nominal interaction with age on BDR, being the strongest interaction observed for the 2-hydroxyglutarate. Results of this metabolite were replicated in Hispanic children with asthma from GACRS \((n = 320)\), with a mean age of 9.1 years. However, these results would not be considered significant after multiple comparison adjustment.

Kelly et al. also conducted a multi-omic study of lung function in Hispanic/Latino children with asthma \((n = 325)\) simultaneously analyzing gene transcripts and metabolites with WGCNA \([49]\). Four transcript modules and five metabolite clusters were found to be related to lung function and seven of them were found to interact among them. Interestingly, one transcriptomic module was associated with BDR and with a lipid metabolomic module, and had as a hub the well-known pediatric-asthma loci ORMDL3, which is involved in sphingolipid biosynthesis \([50][51][52]\). Furthermore, a SNP from ORMDL3 was found to be an eQTL of ORMDL3 and was associated with several lipids included in the metabolomic module. Reinforcing these results, findings were replicated in children with asthma from CAMP \((n = 207)\).

Microbiome

The study of the microbiome involves the analysis of the genetic composition of the microbial communities, known as microbiota. The advances of next-generation sequencing (NGS) techniques have allowed the extension of this type of study by means of targeted sequencing techniques, such as bacterial profiling analyzing the 16S ribosomal RNA gene (16S rRNA), or metagenomic approaches \([53][54]\).

Zhou et al. \([55]\) performed a longitudinal study to identify changes in nasal microbiota related to asthma exacerbations risk despite ICS treatment in European children with mild–moderate persistent asthma \((n = 214)\) as part of a clinical trial. The nasal bacterial microbiome was characterized by means of 16S rRNA sequencing by collecting nasal swabs at the time of well-controlled asthma (randomization) and during the first episode of loss of asthma control. A nasal microbiome dominated by Corynebacterium and Dolosigranulum at randomization was associated with fewer episodes of early loss of asthma control and longer times to develop such episodes. Additionally, during the first loss-of-asthma-control episode, Streptococcus was the most prevalent dominant genus in the nasal microbiome. Moreover, higher bacterial richness and total bacterial load were observed during asthma control loss episode. Furthermore, Corynebacterium was protective for exacerbations requiring OCS use, and the predominance of Moraxella instead of Corynebacterium + Dolosigranulum, was associated with the highest risk of exacerbations. Moreover, Moraxella, Streptococcus, and Haemophilus are pathogens commonly found in the nasal microbiome of asthma patients \([56][57][58]\), and Moraxella has also been associated with a higher risk of asthma exacerbations \([59]\). In contrast, Corynebacterium is the most abundant commensal bacteria in the nasal microbiome from healthy individuals, and asthma patients show decreased relative abundance of this bacteria \([56][60]\).

Conclusions

Omic studies have allowed the identification of novel biomarkers associated with the response to asthma medications in children. However, some findings have not been validated across populations and fail to
fully explain the differences in treatment response in childhood asthma. This is expected since asthma is a complex and multifactorial disease, and different phenotypes may not share the same underlying mechanisms. However, further advances in omic studies will aid to understand how different biological layers are implicated in disease pathogenesis and also how they could interact among them to determine the response to pharmacological therapies.

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**Keywords**

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