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Four childhood atopic dermatitis subtypes identified from trajectory and severity of disease and internally validated in a large UK birth cohort

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SUMMARY

Background—Atopic dermatitis (AD) disease activity and severity is highly variable during childhood. Early attempts to identify subtypes based on disease trajectory have assessed AD presence over time without incorporating severity.

Objective—To identify childhood AD subtypes from symptom severity and trajectories, and determine associations with genetic risk factors, comorbidities and demographic and environmental variables.

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Conflicts of Interest

AC reports personal fees from Novartis, personal fees from Thermo Fisher Scientific, personal fees from Philips, personal fees from Sanofi, personal fees from Stallergenes Greer, outside the submitted work. KA reports consultancy for Target RWE and institutional grant funding from Pfizer, outside the submitted work. ADI has received consultancy fees from AbbVie, Arena Pharmaceuticals, Benevolent AI, Chugai, Dermavant, Genentech, LEO Pharma, Eli Lilly, Menlo Therapeutics, Novartis, Pfizer, Regeneron, Sanofi, and UCB.

All others report no conflicts of interest.

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Methods—We split data from children in the Avon Longitudinal Study of Parents and Children birth cohort into development and validation sets. To identify subtypes, we ran latent class analyses in the development set on AD symptom reports up to age 14. We regressed identified subtypes on non-genetic variables in mutually adjusted, multiply imputed (genetic: unadjusted, complete-case) multinomial regression analyses. We repeated analyses in the validation set and report confirmed results.

Results—11,866 children contributed to analyses. We identified one Unaffected/Rare class (66% of children) and four AD subtypes: Severe-Frequent (4%); Moderate-Frequent (7%); Moderate-Declining (11%); and Mild-Intermittent (12%). Symptom patterns within the first two subtypes appeared more homogeneous than the last two. Filaggrin null mutations (*FLG*), an AD polygenic risk score (PRS), being female, parental AD and comorbid asthma were associated with higher risk for some or all subtypes; *FLG*, AD-PRS and asthma associations were stronger along a subtype gradient arranged by increasing severity and frequency; *FLG* and AD-PRS further differentiated some phenotypes from each other.

Conclusions—Considering severity and AD trajectories leads to four well-defined and recognisable subtypes. The differential associations of risk factors among and between subtypes is novel and requires further research.

INTRODUCTION

Atopic dermatitis (AD) (eczema) is characterised by considerable heterogeneity including clinical morphology, disease severity, age at presentation, disease course, and comorbidities. Many attempts have been made to address this heterogeneity by identifying AD subtypes, which may improve diagnosis, more accurately estimate clinical prognoses, inform management or better predict treatment efficacy or effectiveness¹. Older literature largely subtyped AD into allergic and non-allergic forms; recent evidence suggests that this dichotomisation is not clinically useful² because IgE levels or sensitization status alone do not predict symptom resolution or treatment response³. Recent efforts using longitudinal cohorts have attempted to identify AD subtypes from trajectories of childhood disease activity⁴⁻⁶, or patterns of atopic disease overall⁷⁻¹⁰. However, none of these categorizations account for disease severity, which is the primary predictor of quality of life, sleep disruption, comorbidities including mental health outcomes, and need for systemic treatments^{11,12}. Identifying factors early in the disease course that could differentiate between mild but persistent cases and severe but resolving cases is important both for understanding aetiology and outcomes and for informing patient prognoses and treatment options.

In this paper, we aim to identify AD subtypes based on trajectories of flexural dermatitis in children related to both presence and severity of disease, and quantify their relationships with known and suspected AD risk factors. We note that subtypes based on disease trajectories over time are not the same as directly observed phenotypes¹³. However, as the term 'phenotypes' is commonly used to describe AD subtypes, we use this nomenclature in this paper.

MATERIALS AND METHODS

Study population

We used data from the Avon Longitudinal Study of Parents and Children (ALSPAC)^{14,15}. Pregnant women resident in Avon, UK with expected dates of delivery between 1st April 1991 to 31st December 1992 were invited to take part and 14,541 pregnancies were enrolled initially. Of these, there were 14,676 foetuses, resulting in 14,062 live births and 13,988 children who were alive at 1 year of age, 96% of whom are of white ethnicity. Please note that the study website contains details of all the data that is available through a fully searchable data dictionary and variable search tool¹⁶.

AD phenotype determination

AD presence was collected from questionnaires at eleven ages between 6 and 166 months (13.8 years). Mothers reported whether their children experienced flexural dermatitis (FD) via The International Study of Asthma and Allergies in Childhood (ISAAC)-validated question¹⁷: "Has your child had an itchy, dry skin rash in the joints and creases of his body (e.g., behind the knees, elbows, under the arms) in the past year?" Participants who responded 'yes' were followed up with an assessment of self-rated severity, "How bad was this?" with options: "no problem", "mild", "quite bad" and "very bad". From these two responses we derived a five-category severity variable. Participants who did not complete the questionnaire or did not answer these questions were coded as 'missing'.

Other variables

Family history of atopic diseases, environment exposures and social class were taken from parental questionnaires during pregnancy. Breastfeeding, comorbidities and siblings were taken from questionnaires at various ages (definitions in Supporting Information).

Genetic polymorphisms

Deoxyribonucleic acid (DNA) was obtained from blood samples using standard extraction techniques 18 . In children with genome-wide genotyping data, we created an AD polygenic risk score (PRS) using the 25 single nucleotide polymorphisms (SNPs) previously identified as associated with AD (P<5x10⁻⁰⁸)¹⁹, where dosage genotypes were extracted and weighted by the (risk-increasing) genome-wide association study (GWAS) beta coefficient, summed and standardised to unit standard deviation (1-SD).

In children with filaggrin (*FLG*) loss-of-function genotyping data, null mutations in any of R2447X, S3247X, 2282del4 or R501X were coded as *FLG*-present, and *FLG*-absent was coded otherwise.

Statistical methods

We randomly split the ALSPAC cohort into two equally-sized groups, one (development cohort) for identifying AD phenotypes and associated characteristics, and the other (validation cohort) for internally validating the identified phenotypes. In both groups we excluded individuals with fewer than two of the 11 requested FD reports.

Phenotype identification—We identified phenotypes in the development group using latent class analysis (LCA). FD variables were modelled as joint categorical outcomes in children with and without FD symptoms to allow the LCA to identify a negative control phenotype without AD in addition to AD phenotypes.

We ran models with three to eight classes and compared various model fit statistics (Supporting Information), estimated class sizes, and clinical interpretation (by clinicians with AD management expertise) to determine the best classification. From the best fitting model, we estimated individuals' posterior class probabilities and identified their most likely phenotype as the class with the highest probability. We drew heatmaps for each phenotype showing individuals' symptom severity across ages, with a dendrogram to visualise common patterns.

Association of variables with phenotypes—We explored three methods²⁰ to account for phenotype uncertainty (LCA class probabilities) in regression models (Figure S1). We chose the method that ignores uncertainty and treats phenotypes as known quantities, and report 99% confidence intervals to correct for this method's potential for over-precision.

We regressed sex, family history of AD, asthma, and hayfever, parental education, pet cats and dogs, maternal smoking, breastfeeding (ever and duration), comorbid asthma and older siblings in the household on phenotypes in unadjusted and mutually adjusted multinomial regressions. These factors have been previously associated with AD²¹. In the mutually adjusted model, we excluded 'ever breastfed' to avoid collinearity with breastfeeding duration and additionally adjusted for *FLG* null mutations to investigate independence of family history risk factors. We estimated crude genetic risk by regressing *FLG*, SNPs and the PRS on AD phenotypes in unadjusted multinomial regressions, treating SNP and PRS effects as linear.

Phenotype validation—To assess internal validity, we ran the final LCA model in the validation cohort and visually compared resulting phenotypes with those from the development cohort. We repeated the multinomial regressions and qualitatively compared results with development cohort results.

Missing data handling—We assumed missing FD reports were at random given observed reports and estimated the LCA using full information maximum likelihood (FIML). We checked the solution for sensitivity to departures from this assumption (Supporting Information). We assumed missing non-genetic regression variables were at random given observed variables, social class and comorbid hayfever, and used the multivariate imputation by chained equations²² procedure to impute them. We ran regression models on 10 imputed datasets and combined results using Rubin's rules²³. We performed complete-case genetic analyses, assuming genetic-data missingness was completely at random.

Software—LCAs were run in MPlus v8.2 (Muthén & Muthén, Los Angeles, CA, USA) and all other analyses in Stata v16.0 (StataCorp, College Station, TX, USA) and R v3.6.2 (R Foundation for Statistical Computing, Vienna, Austria).

Ethical approval

Ethical approval for this study was obtained from the ALSPAC Ethics and Law Committee (B2510) and the London School of Hygiene and Tropical Medicine Institutional Review Board (EPNCZK46).

RESULTS

Sample characteristics

13,972 children born as singletons or twins were eligible for inclusion. We randomly allocated half (n=6986) to each of the development and validation cohorts. Within each cohort, 85% had data on FD on two or more questionnaires (development: 5927; validation: 5939) and were included in the analyses. In the development and validation cohorts respectively, just under half were female (48%, 49%); 7.7% (7.4%) carried a *FLG* null mutation and 28% (29%), 19% (18%) and 43% (43%) had at least one parent with a history of AD, asthma and hayfever respectively. All characteristics were balanced between cohorts (Table 1). Reports of FD activity decreased over time. Generally, the activity peaked at 18 months while the lowest activity was observed at age 166 months; similar to the corresponding response rates (92%, 91% at 18 months and 57%, 56% at 166 months) (Table S1).

AD phenotypes from infancy to adolescence

From data in the development cohort, we determined the optimal number of phenotypes from LCA models to be five, based on the Bayesian information criterion (Figure S2) and clinical interpretability (Figure S3). Children were generally clearly classified into phenotypes, with high probability of being in their most likely phenotype (mean 74-90%) and low probabilities of being in the other four (mean 0.1%-10%) (Figure S4, Table S2). There was no evidence the LCA solution was sensitive to missing data patterns and it was consistent across the development and validation cohorts (Figures S5-S6, Tables S2-S3).

Based on reported severity and duration of FD, we labelled the classes: Unaffected/Rare, to which 66% of children were assigned with highest probability; Severe-Frequent AD (4%); Moderate-Frequent AD (7%); Moderate-Declining AD (11%); and Mild-Intermittent AD (12%). Children in a particular phenotype did not always report the same symptoms as each other, but heatmaps in children with complete data showed similar patterns with more consistency among individuals in each Frequent phenotype than among individuals in the Declining and Intermittent phenotypes (Figure S7).

Figure 1 illustrates the distributions of FD severity at each age and how they changed over time for each phenotype. Children in the Severe-Frequent phenotype almost always reported FD and rarely characterised it as 'no problem'. For example, at six months of age, FD was present and rated 'very bad' or 'quite bad' in 29% of the group, rising to 49% at 18 months; although heterogeneity was present within this and other subtypes. Children in the Moderate-Frequent phenotype also almost always reported FD, though they usually rated it less severely than children with Severe-Frequent AD and occasionally rated it no problem. Children with Moderate-Declining AD reported symptoms similar to those with

Moderate-Frequent AD up to age 2.5, but after this age, their likelihood of FD declined. Children with Mild-Intermittent AD more often reported absence of FD than presence, and when present usually rated it mild or no problem. Up to 9% of the Unaffected/Rare group reported FD before age 2.5, mostly mild, but rarely reported it (<=3% of the time) thereafter. Proportions of individuals with missing symptoms are shown in Figure S8.

The distribution of children among phenotypes in the validation cohort was similar to the distribution in the development cohort, except that relatively fewer children were assigned to the Mild-Intermittent phenotype (8% vs 14%) and more to Unaffected/Rare (66% vs 62%).

Variables associated with phenotypes

Table 2 shows the distributions of clinical, demographic and genetic variables used in the regression analyses by most likely phenotype.

We used the Unaffected/Rare phenotype as the reference group in multiple imputation analyses (Table S4). The unadjusted and adjusted estimates were similar, except for parental asthma, hayfever and education: each unadjusted estimate showed a higher risk for all but Mild-Intermittent AD, which attenuated after adjustment. The adjusted estimates in the development and validation cohorts were also similar (Figure 2).

After adjusting for the other variables and *FLG*, we found that comorbid asthma, parental AD and sex were associated with AD phenotype (all p<0.0001). Children with asthma were three times more likely to be in the Severe-Frequent group (relative risk ratio (RRR) 3.06 99% confidence interval (CI) [1.80, 5.20]), twice as likely to be Moderate-Frequent (RRR 1.98 [1.41, 2.79]); and 1.41 (1.02, 1.94) and 1.44 (1.07, 1.95) times as likely to be Moderate-Declining and Mild-Intermittent, respectively. There was strong evidence (p=0.0001) for this differential association with each phenotype. Children with a family history of AD had twice the risk of Severe-Frequent (RRR 1.97, 1.32, 2.94) or Moderate-Frequent (RRR 1.91 [1.40, 2.62]) disease, 1.61 (1.24, 2.09) times the risk of Moderate-Declining disease and 1.39 (1.07, 1.81) times the risk of Mild-Intermittent disease, but there was no evidence that these associations were truly differential (p=0.0987). Girls were more likely than boys to have Mild-Intermittent AD (RRR 1.67 (1.34, 2.08)). There was no evidence that the other Table 2 variables were independently associated with AD phenotypes.

The patterns of associations above were replicated in the validation cohort with similar estimates and confidence intervals, except that differential parental AD RRRs were detectable across phenotypes (p=0.0039). Additionally, in the validation cohort there was borderline evidence (p=0.0063) that maternal smoking was associated with phenotypes after adjusting for other variables; there was similar evidence (p=0.0135) for the same association in the development cohort. Children whose mothers smoked were less likely to have Moderate-Frequent AD in the development cohort and less likely to have Moderate-Declining AD in the validation cohort; the other associations were inconclusive.

Genetic phenotype associates

Of children included in this study, 8773 (74%) had *FLG* genotype data and 7782 (66%) had available genome-wide genotype data.

FLG loss-of-function mutations—FLG was strongly associated with AD phenotypes (p=4.26x10⁻³²), with strong evidence of higher risks for more severe and frequently-active phenotypes (p=1.23x10⁻¹⁰). Children with FLG mutations were more than four times more likely to have Severe-Frequent AD (RRR 4.32 [3.07, 6.07]), more than twice as likely to have Moderate-Frequent AD (RRR 2.41 [1.81, 3.22]), 1.80 (1.39, 2.34) times more likely to have Moderate-Declining AD and 1.52 (1.13, 2.04) times more likely to have Mild-Intermittent AD than children without FLG mutations (Table 3).

Standardised Genetic Score—The PRS was strongly associated with AD phenotypes $(p=1.22\times10^{-28})$, with strong evidence of higher risks for more severe and frequently-active phenotypes $(p=8.60\times10^{-11})$. A score increase of 1-SD was associated with a relative risk increase of: 1.67 (1.45, 1.93) for Severe-Frequent; 1.37 (1.23, 1.52) for Moderate-Frequent; 1.17 (1.07, 1.27) for Moderate-Declining; and 1.13 (1.02, 1.24) for Mild-Intermittent AD. Removing rs11205006 from the score, which tags the *FLG* locus, and the remainder of the Chromosome 1 SNPs made little difference to these associations (Table S5).

SNPs—Eight SNPs on Chromosomes 1, 6, 11, 14 and 19 were associated with AD phenotypes (p= 8.63×10^{-10} - 1.19×10^{-02} ; Table 3). Of these, five were associated with Severe-Frequent AD (RRR range 1.31-2.99); six with Moderate-Frequent (RRR range 1.17-2.07); and one with Moderate-Declining (RRR 1.96). There was no evidence that any single SNP was associated with Mild-Intermittent AD.

Association differences between AD phenotypes

We explored whether any of the variables we investigated could distinguish between the four AD phenotypes. In six *post-hoc* Bonferroni-corrected pairwise comparisons from the multinomial regression models, we found evidence that the relative risk associated with asthma, sex, *FLG* null mutations and the AD-PRS differed among some phenotype pairs (Table S6). In particular, the AD-PRS and FLG were able to discriminate between four of six pairs showing a significantly higher relative risk in the more severe and/or frequently affected phenotypes. The remaining comparisons were inconclusive, generally producing confidence intervals consistent with important effects in both directions.

DISCUSSION

We identified four phenotypes of AD based on disease severity and trajectories during childhood: Severe-Frequent, Moderate-Frequent, Moderate-Declining and Mild-Intermittent disease. Providers and patients might be reassured that in all subgroups, the probability of reporting 'quite bad' or 'very bad' symptoms declined with age (with evidence of within group heterogeneity). Associations with some established risk factors (*FLG* null mutations, AD-PRS, family history of AD) and comorbidities (asthma) were present in all phenotypes with evidence of a gradient of association from more severe to less severe phenotypes. AD-PRS and *FLG* mutations were further able to differentiate many of our phenotypes from each other, highlighting the need for increased understanding of drivers of disease severity/activity throughout life in people with AD, including the role of gene-environment interactions.

Our findings add novel information about patterns of disease severity over time, and are consistent with prior cross-sectional studies on disease severity and longitudinal studies focused on disease activity alone. While disease activity (i.e. the presence of any symptoms at a particular age) and disease severity (i.e. the impact of those symptoms) are sometimes correlated, they are separate concepts, and it is important to differentiate them²⁴. For example, a small patch of mild eczema that persists for years may be quite different from extensive severe disease that quickly resolves.

Paternoster et al⁵ conducted a previous study in ALSPAC with external validation²⁵, Hu et al⁶ examined a multi-ethnic population up to age 10, and Roduit et al⁴ examined an European population up to age 6. All reported phenotypes described onset timing and persistence during childhood, and between them there were some commonly described phenotypes (early-persistent, early-transient, late-onset). Hu et al concluded that established risk factors have little capacity to differentiate eczema phenotypes. By additionally incorporating severity, the most clinically relevant aspect of AD, we have been able to show that genetic factors, in particular, have potential to differentiate between phenotypes.

Our study has several strengths. We used a well-characterised, population-based cohort, hence, our results are generalisable to similar settings and populations. Our large sample (n=11,866) offered precision; and our phenotypes were internally reproducible, lending confidence to our findings. We applied robust statistical methods to minimise imprecision and bias in our regression estimates.

Our study also has limitations. We used a measure of self-reported AD, which has not been directly validated in ALSPAC, but has been shown to be a reasonable predictor in other contexts, including validation by physical exam in the UK²⁶ and physician assessment in a multi-centre US study²⁷. Missing data could have introduced bias into our estimates, although we tried to minimise this risk with FIML estimation and multiple imputation techniques. There is uncertainty in phenotype assignment that could affect regression results, however, we compared several methods of accounting for it and found little difference in our results, suggesting insensitivity to this issue. Finally, there are few data sources recording AD severity, and we were unable to find a large cohort with similarly characterised children to externally validate our findings, though we did attempt this using a small cohort (Supporting Information).

The LCA method involves probabilistic classification into subgroups and individual disease trajectories can vary within subgroups. For example, on average, active disease becomes less likely over time in the Moderate-Declining group, but an individual classified into this phenotype may still have active FD in late childhood. Future work may consider how early aggressive treatment for severe AD might modify the natural history²⁸.

Our study adds important information about patterns of AD severity over time. The identified phenotypes were stable across sensitivity analyses and had strong genetic associations, providing support for using them in future work toward a better aetiological understanding of AD, including the role of environmental risk factors throughout the life

course, and toward better clinical trial design. Additional research is underway to determine the utility of our phenotypes for prognosis and to inform priorities for early intervention.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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What's already known about this topic?

• Atopic dermatitis (AD) is a heterogeneous condition in terms of both disease activity and severity.

 Childhood AD phenotypes in previous studies have focused only on disease activity.

What does this study add?

- We incorporate disease severity over time to derive four clinically recognisable AD phenotypes using data-driven methods.
- Disease severity improved over time in all phenotypes (even in those with high probability of activity in late childhood and adolescence).
- Several established risk factors, including genetics associates, were associated with our proposed phenotypes, with most factors more strongly associated with phenotypes reporting the worst symptoms. Fewer factors differentiated between more frequent and declining/intermittent phenotypes.

Phenotypes of atopic dermatitis

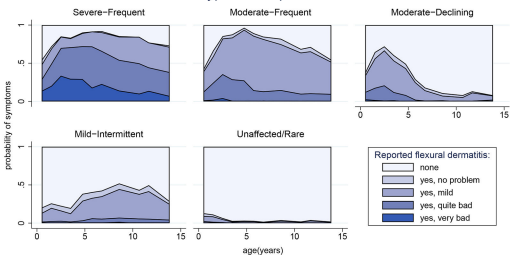
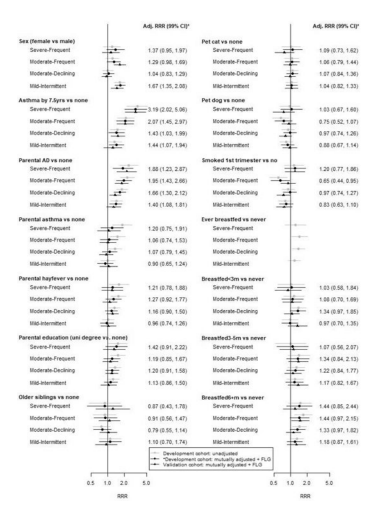


Figure 1: Atopic dermatitis phenotypes

Five phenotypes of atopic dermatitis (including an Unaffected/Rare phenotype without atopic dermatitis) identified by latent class analysis on 5927 ALSPAC children whose mothers responded to at least two questionnaires between the ages of 6 months and 14 years. Probabilities of flexural dermatitis and its severity are shown by age for children assigned to each phenotype (missing reports not indicated): Severe-Frequent, n=230 (3.9%); Moderate-Frequent, n=408 (6.9%); Moderate-Declining, n=676 (11%); Mild-Intermittent, n=684 (12%); Unaffected/Rare, n=3929 (66%).



 $\label{thm:continuous} \textbf{Figure 2: Associations of demographic and clinical characteristics with atopic dermatitis phenotypes } \\$

Unadjusted and mutually adjusted phenotype associations in the development cohort (grey and black circles), and adjusted associations in the validation cohort (black triangles). All adjusted associations include additional adjustment for *FLG* null mutations. Stated effects and confidence intervals are for adjusted associations in the development cohort. Adj=adjusted; RRR=relative risk ratio; CI=confidence interval.

Table 1:

ALSPAC study characteristics

	Development cohort (n=5927)	Validation cohort (n=5939)
Sex		
Male	3099 (52%)	3031 (51%)
Female	2828 (48%)	2908 (49%)
Social class *	'	
I	693 (12%)	677 (11%)
П	2031 (34%)	1999 (34%)
III(n)	1203 (20%)	1219 (21%)
III(m)	387 (6.5%)	414 (7%)
IV	113 (1.9%)	125 (2.1%)
V	18 (0.3%)	7 (0.1%)
Missing	1482 (25%)	1498 (25%)
At least one parent with university degree		
No	4393 (74%)	4420 (74%)
Yes	1288 (22%)	1238 (21%)
Missing	246 (4.1%)	281 (4.7%)
Parental AD		
No	2563 (43%)	2557 (43%)
Yes	1678 (28%)	1700 (29%)
Missing	1686 (28%)	1682 (28%)
Parental asthma		
No	2967 (50%)	3041 (51%)
Yes	1123 (19%)	1059 (18%)
Missing	1837 (31%)	1839 (31%)
Parental hayfever		
No	1826 (31%)	1812 (31%)
Yes	2534 (43%)	2549 (43%)
Missing	1567 (26%)	1578 (27%)
Any older siblings		
No	433 (7.3%)	454 (7.6%)
Yes	2105 (35%)	1962 (33%)
Missing	3389 (57%)	3523 (59%)
Pet cat during pregnancy		
No	4011 (68%)	3965 (67%)
Yes	1723 (29%)	1778 (30%)
Missing	193 (3.2%)	196 (3.3%)
Pet dog during pregnancy		

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		Development cohort (n=5927)	Validation cohort (n=5939)
	No	4353 (73%)	4358 (73%)
	Yes	1381 (23%)	1385 (23%)
	Missing	193 (3.2%)	196 (3.3%)
Mum smoked during	g first 3m of pregnancy		
	No	4458 (75%)	4408 (74%)
	Yes	1307 (22%)	1361 (23%)
	Missing	162 (2.7%)	170 (2.9%)
Breastfeeding			
	Never	1409 (24%)	1349 (23%)
	<3mths	1214 (20%)	1240 (21%)
	3-5mths	855 (14%)	908 (15%)
	6mths+	1857 (31%)	1802 (30%)
	missing	592 (10%)	640 (11%)
Asthma by 7.5 yrs			
	No	3242 (55%)	3188 (54%)
	Yes	816 (14%)	829 (14%)
	Missing	1869 (31%)	1922 (32%)
Hayfever by 10.5 year	ars		
	No	2877 (48%)	2856 (48%)
	Yes	784 (13%)	732 (12%)
	Missing	2266 (38%)	2351 (40%)
FLG null mutation			
	No	3985 (67%)	3891 (66%)
	Yes	456 (7.7%)	441 (7.4%)
	Missing	1486 (25%)	1607 (27%)
Polygenic risk score	(standardised)		
	Median [IQR]	-0.13 [-0.78, 0.63]	-0.10 [-0.74, 0.62]
	Missing	1971 (33%)	2113 (36%)

^{*} I Professional occupations, II Managerial and Technical occupations, III(n) Skilled occupations-non-manual, III(m) Skilled occupations-manual, IV Partly-skilled occupations, V Unskilled occupations.

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 Table 2:

 ALSPAC development cohort characteristics across each 'most likely' atopic dermatitis phenotype

	Severe- Frequent	Moderate- Frequent	Moderate- Declining	Mild- Intermittent	Unaffected /Rare
Phenotype N (%)	230 (3.9%)	408 (6.9%)	676 (11%)	684 (12%)	3929 (66%)
Sex	,				
Male	115 (3.7%)	202 (6.5%)	366 (12%)	287 (9.3%)	2129 (69%)
Female	115 (4.1%)	206 (7.3%)	310 (11%)	397 (14%)	1800 (64%)
At least one parent w	rith university degree			,	
No	158 (3.6%)	286 (6.5%)	480 (11%)	502 (11%)	2967 (68%)
Yes	63 (4.9%)	113 (8.8%)	170 (13%)	164 (13%)	778 (60%)
Missing	9 (3.7%)	9 (3.7%)	26 (11%)	18 (7.3%)	184 (75%)
Parental AD	,	,		,	,
No	73 (2.8%)	138 (5.4%)	278 (11%)	296 (12%)	1778 (69%)
Yes	95 (5.7%)	170 (10%)	249 (15%)	218 (13%)	946 (56%)
Missing	62 (3.7%)	100 (5.9%)	149 (8.8%)	170 (10%)	1205 (71%)
Parental asthma	,				,
No	99 (3.3%)	192 (6.5%)	344 (12%)	364 (12%)	1968 (66%)
Yes	61 (5.4%)	99 (8.8%)	155 (14%)	124 (11%)	684 (61%)
Missing	70 (3.8%)	117 (6.4%)	177 (9.6%)	196 (11%)	1277 (70%)
Parental hayfever					
No	56 (3.1%)	106 (5.8%)	198 (11%)	232 (13%)	1234 (68%)
Yes	122 (4.8%)	207 (8.2%)	325 (13%)	287 (11%)	1593 (63%)
Missing	52 (3.3%)	95 (6.1%)	153 (9.8%)	165 (11%)	1102 (70%)
Any older siblings					
No	17 (3.9%)	38 (8.8%)	62 (14%)	49 (11%)	267 (62%)
Yes	86 (4.1%)	168 (8%)	248 (12%)	285 (14%)	1318 (63%)
Missing	127 (3.7%)	202 (6%)	366 (11%)	350 (10%)	2344 (69%)
Pet cat during pregna	ancy				
No	156 (3.9%)	277 (6.9%)	453 (11%)	466 (12%)	2659 (66%)
Yes	67 (3.9%)	123 (7.1%)	205 (12%)	205 (12%)	1123 (65%)
Missing	7 (3.6%)	8 (4.1%)	18 (9.3%)	13 (6.7%)	147 (76%)
Pet dog during pregn	ancy				
No	168 (3.9%)	327 (7.5%)	505 (12%)	524 (12%)	2829 (65%)
Yes	55 (4%)	73 (5.3%)	153 (11%)	147 (11%)	953 (69%)
Missing	7 (3.6%)	8 (4.1%)	18 (9.3%)	13 (6.7%)	147 (76%)
	first 3 months of pregnan				
No	164 (3.7%)	337 (7.6%)	517 (12%)	535 (12%)	2905 (65%)
Yes	59 (4.5%)	63 (4.8%)	146 (11%)	131 (10%)	908 (69%)
Missing	7 (4.3%)	8 (4.9%)	13 (8%)	18 (11%)	116 (72%)

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	Severe- Frequent	Moderate- Frequent	Moderate- Declining	Mild- Intermittent	Unaffected /Rare
Breastfeeding					
Never	49 (3.5%)	81 (5.7%)	139 (9.9%)	160 (11%)	980 (70%
<3mths	40 (3.3%)	76 (6.3%)	160 (13%)	128 (11%)	810 (67%)
3-5mths	27 (3.2%)	64 (7.5%)	98 (11%)	109 (13%)	557 (65%)
6mths+	86 (4.6%)	157 (8.5%)	231 (12%)	238 (13%)	1145 (62%)
Missing	28 (4.7%)	30 (5.1%)	48 (8.1%)	49 (8.3%)	437 (74%)
Asthma by 7.5 years					
No	96 (3%)	211 (6.5%)	360 (11%)	417 (13%)	2158 (67%)
Yes	62 (7.6%)	90 (11%)	106 (13%)	119 (15%)	439 (54%)
Missing	72 (3.9%)	107 (5.7%)	210 (11%)	148 (7.9%)	1332 (71%)
FLG null mutation					
No	121 (3%)	261 (6.5%)	461 (12%)	481 (12%)	2661 (67%)
Yes	55 (12%)	55 (12%)	66 (14%)	58 (13%)	222 (49%)
Missing	54 (3.6%)	92 (6.2%)	149 (10%)	145 (9.8%)	1046 (70%)
Polygenic risk score (standa	ardised)				
Median [IQR]	0.39 [-0.38, 1.47]	0.18 [-0.73, 0.98]	-0.03 [-0.74, 0.76]	-0.02 [-0.73, 0.64]	-0.22 [-0.82, 0.54]
Missing*	73 (32%)	125 (31%)	200 (30%)	189 (28%)	1384 (35%

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^{*} missing polygenic risk scores are tabulated within phenotype, while missingness in other variables is tabulated across phenotypes.

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Table 3:

Associations between genetic profiles and atopic dermatitis phenotypes in ALSPAC children

				Severe-Frequent* N=305	Moderate- Frequent* N=634	Moderate- Declining* N=1008	Mild-Intermittent* N=845	
	Chr.	Nearest gene	Global p- value [†]	RRR (99% CI)	RRR (99% CI)	RRR (99% CI)	RRR (99% CI)	Equality p- value [‡]
FLG null mutation			4.26E-32	4.32 (3.07, 6.07)	2.41 (1.81, 3.22)	1.80 (1.39, 2.34)	1.52 (1.13, 2.04)	1.23E-10
Polygenic risk score	e		1.22E-28	1.67 (1.45, 1.93)	1.37 (1.23, 1.52)	1.17 (1.07, 1.27)	1.13 (1.02, 1.24)	8.60E-11
Individual SNP associations	ociations							
rs11205006_a	1	LCE5A, FLG-AS1	9.94E-07	1.55 (1.24, 1.93)	1.22 (1.03, 1.44)	1.11 (0.97, 1.28)	1.10 (0.95, 1.28)	3.20E-03
rs2228145_c	1	IL6R	3.69E-04	1.31 (1.06, 1.63)	1.15 (0.99, 1.34)	0.95 (0.83, 1.08)	0.93 (0.81, 1.07)	1.49E-04
rs61813875_g	1	LCE3E, CRCT1	8.63E-10	2.99 (1.79, 4.98)	2.07 (1.34, 3.19)	1.96 (1.36, 2.84)	1.33 (0.84, 2.09)	9.82E-03
$\rm rs1057258_c$	2	INPP5D	5.81E-02	1.21 (0.90, 1.63)	0.99 (0.81, 1.21)	0.97 (0.82, 1.14)	0.86 (0.72, 1.01)	4.81E-02
rs112111458_a	2	CD207	6.68E-02	1.18 (0.85, 1.65)	1.00 (0.80, 1.25)	1.17 (0.96, 1.42)	0.91 (0.75, 1.10)	4.42E-02
rs13015714_g	2	IL18R1	5.40E-02	1.07 (0.83, 1.38)	1.23 (1.03, 1.47)	1.06 (0.92, 1.24)	1.02 (0.87, 1.20)	1.74E-01
rs17389644_a	4	IL21	9.94E-02	1.17 (0.91, 1.51)	1.07 (0.89, 1.29)	1.15 (0.99, 1.33)	1.02 (0.87, 1.21)	4.36E-01
$rs10214237_t$	5	IL7R	3.01E-01	0.96 (0.75, 1.21)	1.06 (0.89, 1.26)	1.01 (0.88, 1.17)	1.13 (0.96, 1.32)	3.66E-01
rs2897442_c	5	KIF3A	4.78E-02	1.29 (1.02, 1.63)	1.11 (0.93, 1.32)	0.99 (0.86, 1.15)	0.99 (0.85, 1.16)	4.02E-02
rs12153855_t	9	TNXB	8.75E-03	1.04 (0.72, 1.50)	1.42 (1.06, 1.91)	1.07 (0.86, 1.33)	1.22 (0.96, 1.56)	1.32E-01
rs6473227_c	8	ZBTB10	2.49E-01	1.12 (0.90, 1.40)	1.09 (0.93, 1.28)	1.07 (0.94, 1.22)	1.06 (0.93, 1.22)	9.47E-01
$rs10995251_c$	10	ZNF365	5.82E-01	0.99 (0.79, 1.23)	0.99 (0.84, 1.16)	1.08 (0.95, 1.24)	1.03 (0.89, 1.18)	5.87E-01
rs12295535_t	11	PRR5L	8.22E-03	1.55 (0.92, 2.61)	1.21 (0.80, 1.84)	1.37 (0.99, 1.91)	0.78 (0.50, 1.22)	1.83E-02
rs479844_g	11	OVOL1	1.17E-05	1.51 (1.21, 1.90)	1.17 (1.00, 1.37)	1.06 (0.93, 1.21)	1.06 (0.92, 1.22)	1.34E-03
$rs7127307_t$	11	ETS1	5.28E-01	1.05 (0.84, 1.30)	1.07 (0.92, 1.25)	1.00 (0.88, 1.13)	1.07 (0.94, 1.23)	6.39E-01
rs7927894_t	11	C11orf30	2.48E-02	1.21 (0.98, 1.50)	1.09 (0.94, 1.28)	0.96 (0.84, 1.09)	1.10 (0.96, 1.26)	3.74E-02
rs2227483_t	12	IL22	9.80E-01	1.03 (0.83, 1.29)	1.01 (0.86, 1.18)	1.00 (0.88, 1.14)	1.03 (0.90, 1.18)	9.75E-01
$rs2143950_t$	14	PPP2R3C	1.19E-02	1.24 (0.95, 1.62)	1.26 (1.04, 1.53)	1.01 (0.86, 1.20)	1.06 (0.89, 1.27)	5.74E-02
$\rm rs7146581_c$	14	TRAF3	1.83E-01	1.09 (0.84, 1.42)	1.16 (0.96, 1.40)	1.09 (0.94, 1.27)	1.04 (0.88, 1.22)	6.78E-01
rs2041733_t	16	CLEC16A	4.41E-01	0.98 (0.79, 1.22)	1.12 (0.96, 1.30)	1.00 (0.88, 1.14)	1.03 (0.90, 1.18)	4.20E-01
rs11657987_t	17	PGS1	9.36E-02	1.22 (0.99, 1.51)	1.09 (0.94, 1.27)	1.02 (0.90, 1.15)	1.06 (0.92, 1.21)	2.44E-01

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				Severe-Frequent* Moderate- N=305 Frequent* N=634		Moderate- Declining* N=1008	Mild-Intermittent* N=845	
	Chr.	Chr. Nearest gene	Global p-value †	Global p- RRR (99% CI) RRR (99% CI) RRR (99% CI) RRR (99% CI) value $\mathring{\it f}$	RRR (99% CI)	RRR (99% CI)		Equality p- value [‡]
rs16948048_g	17	ZNF652	7.69E-02	1.18 (0.95, 1.46)	0.92 (0.79, 1.08)	0.98 (0.86, 1.11)	7.69E-02 1.18 (0.95, 1.46) 0.92 (0.79, 1.08) 0.98 (0.86, 1.11) 0.92 (0.80, 1.06)	5.50E-02
rs17881320_t	17	STAT3	2.25E-01	1.30 (0.91, 1.86)	1.12 (0.85, 1.47)	1.13 (0.90, 1.41)	2.25E-01 1.30 (0.91, 1.86) 1.12 (0.85, 1.47) 1.13 (0.90, 1.41) 1.09 (0.86, 1.39)	7.37E-01
rs2164983_a	19	ACTL9	1.26E-05	1.26E-05 1.34 (1.02, 1.77) 1.40 (1.15, 1.70) 1.16 (0.98, 1.37) 1.18 (0.99, 1.42)	1.40 (1.15, 1.70)	1.16 (0.98, 1.37)	1.18 (0.99, 1.42)	1.49E-01
rs6010620_g	20	RTEL1	9.39E-02	9.39E-02 1.14 (0.88, 1.47) 1.17 (0.97, 1.41) 1.05 (0.90, 1.21) 1.11 (0.94, 1.30)	1.17 (0.97, 1.41)	1.05 (0.90, 1.21)	1.11 (0.94, 1.30)	6.05E-01

compared with the Unaffected/Rare population, n=4990

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[†] null hypothesis: RRR1=RRR2=RRR3=RRR4. FLG=filaggrin gene; SNP=single nucleotide polymorphism; Chr=chromosome; RRR=relative risk ratio; CI=confidence interval.