Insights into the genetic architecture of the human face

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The human face is complex and multipartite, and characterization of its genetic architecture remains challenging. Using a multivariate genome-wide association study meta-analysis of 8,246 European individuals, we identified 203 genome-wide-significant signals (120 also study-wide significant) associated with normal-range facial variation. Follow-up analyses indicate that the regions surrounding these signals are enriched for enhancer activity in cranial neural crest cells and craniofacial tissues, several regions harbor multiple signals with associations to different facial phenotypes, and there is evidence for potential coordinated actions of variants. In summary, our analyses provide insights into the understanding of how complex morphological traits are shaped by both individual and coordinated genetic actions.
Furthermore, we reveal interactions between variants at different loci affecting similar aspects of facial shape variation, identifying gene sets that work in concert to build human faces. With this work, we not only push forward our understanding of human facial genetics, but also illustrate the potential for researchers to confront Atchley and Hall’s problem, by intensively characterizing complex morphological variation and using advanced methods to identify factors involved in the developmental choreography of complex morphological structures.

Results

Multivariate phenotyping and meta-analysis framework. To study facial variation at both global and local scales, we start with a set of three-dimensional (3D) facial surface scans, upon which we map a dense mesh of 7,160 homologous vertices\textsuperscript{41}. We then apply a data-driven facial segmentation approach, defined by grouping vertices that are strongly correlated using hierarchical spectral clustering\textsuperscript{42,43}. The configurations of each of the resulting 63 segments are then subjected independently to a Generalized Procrustes analysis, after which principal component analysis (PCA) is performed in conjunction with parallel analysis to capture the major phenotypic variation in each facial segment\textsuperscript{44,45} (Extended Data Fig. 1). The number of principal components (PCs) kept at this stage of the analysis ranged from 7 to 70, with segments containing large numbers of quasi-landmarks generally requiring more PCs to describe the variation in that segment. The inherent shape variability in each segment also plays a role in the number of PCs retained by parallel analysis, with more variable segments retaining more PCs. For example, although segments 5 and 25 contain similar numbers of quasi-landmarks, because the variability of the nose (segment 5) is generally greater than that of the lower cheeks (segment 25), the parallel analysis for segment 5 retained 32 PCs while for segment 25 it retained only 20 PCs (Extended Data Fig. 1b).

We then tested for genetic association between the facial PCs and 7,417,619 SNPs by using a data-driven approach (Extended Data Fig. 2). Within each segment, instead of a priori selecting the PCs of interest, or treating each of the 63 segments as a single ‘trait’, we use canonical correlation analysis (CCA) to first identify the linear combination of components in each segment maximally correlated with the SNP being tested in the identification cohort. We call this multivariate combination of PCs the ‘trait’. Thus, each SNP is associated (although not always with significance) with its own ‘trait’ in each segment. Subsequently, the verification cohort is projected onto each of these traits, creating univariate ‘phenotype’ variables that are tested for genotype–phenotype associations by using linear regression. The projection ensures that the shape variation tested in the verification step is equivalent to the ‘trait’ used in the identification step. The identification and verification P values are then meta-analyzed using Stouffer’s method\textsuperscript{46,47}. The whole process is then repeated, switching the dataset used for identification and verification, thereby resulting in 126 meta-analysis P values and traits (63 segments × 2 meta-analysis tracks) for each SNP. Further details are available in the Methods and Supplementary Notes 1 and 2.

Sharing of genome-wide signals between facial segments. We first assessed the degree to which variation in each facial segment shares the same patterns of association across the genome by computing the linkage disequilibrium score correlation (LDSC) based on genome-wide-association P values for each pair of facial segments\textsuperscript{48,49}. This $63 \times 63$ matrix of correlations was visualized on top of the facial segmentation hierarchy to assess between-segment correlations within and between facial quadrants (Extended Data Fig. 3), although it is important to note that these LDSCs should not be considered ‘genetic correlations’ in the typical way of a univariate trait, since the z-scores used are unsigned. The LDSCs were highest between segments of the same facial quadrant (that is, lips, nose, lower face, upper face), validating the hierarchical clustering used to initially define the segments (Extended Data Fig. 3b). Average-linkage hierarchical clustering of the facial segments based on the correlation values gave rise to four main clusters, each corresponding primarily to segments from the same quadrant (Extended Data Fig. 4). Despite substantial within-quadrant similarity, there were notable correlations between groups of segments from different quadrants (Extended Data Fig. 3a). Some of these specific correlations reflect close physical proximity of the segments in different quadrants (for example, segments 12 and 33), but some correlations seem to reflect the shared embryological origins of groups of segments. Specifically, segments representing the nose (quadrant II) and upper face (quadrant IV) cluster together, and most segments representing the lips (quadrant I) and lower face (quadrant III) cluster together (Extended Data Fig. 4). Quadrants II and IV together approximate the frontalonal prominence, which appears earlier in development than the mandibular and maxillary prominences, which are approximated by quadrants I and III, respectively\textsuperscript{50}.

Genome-wide-association meta-analysis. In total, we identified 17,612 SNPs with $P$ values ($P_{\text{Meta-US}}$ and/or $P_{\text{Meta-UK}}$) lower than the genome-wide threshold ($P \leq 5 \times 10^{-8}$). Of these, 11,398 SNPs also passed the study-wide-significance threshold ($P \leq 6.96 \times 10^{-10}$) (Supplementary Fig. 1). For each peak passing the genome-wide threshold, we designated the SNP with the lowest $P$ value across all facial segments as the ‘lead SNP’, refining our results to 218 genome-wide-significant lead SNPs. Of these, 203 SNPs showed consistent genetic effects on the trait identified in the US- and UK-driven meta-analyses in the facial segment with the lowest $P$ value for that SNP (Fig. 1 and Supplementary Table 3), and 120 of these were also below study-wide significance. Visual representations of the LocusZoom\textsuperscript{51} and effect plots for each of the 203 genome-wide-significant SNPs are available in the FigShare repository\textsuperscript{52}.

The global-to-local approach means that we often identified associations between a single SNP and variation in many facial segments. In this article, we focus primarily on the segment in which the SNP had its lowest $P$ value (the ‘Best segment’) and provide information on which in meta-analysis track (Meta-US or Meta-UK) the SNP reached this significance level (the ‘Best meta-analysis track’). Thus, throughout the rest of the article, the reported $P$ values for each SNP will be in the format of $P_{\text{best-track}}$ (Best segment) = value. By plotting the strongest association results for each segment (Fig. 1, left), segments 1 and 2 are visibly the ‘Best segment’ for most SNPs, with $n = 20$ SNPs reaching lowest significance in the full face (segment 1) in the US-driven meta-analysis ($n = 15$ for Meta-UK) and $n = 19$ SNPs reaching lowest significance in segment 2 in the US-driven meta-analysis ($n = 18$ for Meta-UK).

Genes near lead SNPs are enriched for both craniofacial and limb development. In a GREAT\textsuperscript{53} analysis of the regions surrounding the 203 genome-wide-significant lead SNPs, the top ten terms (based on lowest binomial $P$ values) in the mouse phenotype, human phenotype and gene ontology (GO) biological processes categories are all highly relevant to craniofacial shape and overall morphology (Extended Data Fig. 5a), with the top human phenotype being oral clefing. A FUMA\textsuperscript{54} analysis of the same regions highlighted genes overlapping several pathways related to abnormal cellular maintenance and also included pathways highly relevant for morphological development, like the Wnt, Hedgehog and TGFβ signaling pathways (Extended Data Fig. 5b).

Facial GWAS peaks are enriched for enhancers specific to cell types across the timeline of facial development. To assess the likelihood of cell types and developmental timepoints in which our GWAS regions are active, we compiled H3K27ac ChiP–seq signals—detecting a
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Fig. 1 | Overall results of US-driven and UK-driven meta-analyses. On the left, numbered blocks representing the 63 facial segments arranged and colored according to quadrant (I, orange; II, red; III, light blue; IV, dark blue), the full face (white) and segments 2 (light orange) and 3 (ice blue). The histogram arranged on the left side represents the number of genome-wide-significant lead SNPs reaching their lowest P value in each segment, with each rectangle representing one SNP. The US-driven meta-analysis results are on the outside of the circle and the UK-driven meta-analysis results are on the inside of the circle. In the center, the global-to-local facial segmentation of all 3D images included in this analysis, obtained using hierarchical spectral clustering, are colored to match with the quadrants on the left. On the right, a Miami plot of the US-driven meta-analysis P values on the outside and the UK-driven meta-analysis P values on the inside, with chromosomes colored and labeled. Values plotted are the result of Stouffer’s meta-analysis of one-sided right-tailed identification and verification P values, detailed in the Methods, and are –log10 scaled (range, 0–80). The red line represents the genome-wide-significance threshold (P = 5 × 10⁻⁵) and the black line represents the study-wide threshold (P = 6.96 × 10⁻¹⁰). Created using Circos v.0.69-8 (ref. 53).

Marker of the promoters of transcriptionally active genes and active distal enhancers— from approximately 100 different cell types and tissues, including cranial neural crest cells (CNCCs), fetal and adult osteoblasts and mesenchymal stem cell-derived chondrocytes, as well as dissected embryonic craniofacial tissues (Carnegie stages 13–20). Both CNCCs and craniofacial tissues showed the highest H3K27ac signals in the vicinity of the 203 genome-wide-significant lead SNPs, whereas no H3K27ac signal was observed for 203 random SNPs matched for allele frequency and distance to the nearest gene (Fig. 2a). The difference in H3K27ac signal between the 203 genome-wide-significant lead and random SNPs was significant based on a two-sided Wilcoxon rank-sum test for many cell types and tissues, with CNCCs and embryonic craniofacial tissues having the greatest median differences (Extended Data Fig. 6 and Supplementary Table 4).

To distinguish enrichment between coding and noncoding elements, we examined chromatin signals in CNCCs and embryonic craniofacial tissues in more detail, using ChIP–seq data on additional chromatin marks and transcription factors. In the CNCCs, candidate regulatory regions in the vicinity of the 203 genome-wide-significant lead SNPs were enriched significantly for strong and intermediate enhancers and depleted in weak promoters (Fig. 2b). In embryonic craniofacial tissue, all developmental stages sampled were significantly enriched for the chromHMM states of...
active enhancers, active enhancer flanks and weak enhancers, and depleted in quiescent/low and heterochromatin states (Fig. 2c).

Cell-type-specific activity patterns were used to further subdivide the 203 genome-wide-significant lead SNPs by using k-means clustering of H3K27ac signals (Fig. 3). As expected, many lead SNPs showed specific activity for CNCCs and craniofacial tissue (for example, cluster 5), representing activity at an early time point in development. Interestingly, however, some SNPs showed preferential activity for either CNCCs or craniofacial tissue (for example, clusters 1 and 2). Greater specificity for CNCCs could arise because CNCCs constitute a relatively small proportion of the cells present in craniofacial tissue at Carnegie stages 13–20, while greater specificity for craniofacial tissue could be due to activity in further differentiated cell types of the face.

**Known and new loci.** We identified 89 genome-wide-significant (66 also study-wide-significant) peaks that overlap with the results of prior association studies of normal-range facial phenotypes. Of these, 29 genome-wide-significant (20 also study-wide-significant) peaks were reported by studies with overlapping samples as this study and 60 genome-wide-significant (46 also study-wide-significant) peaks were previously reported by studies with completely non-overlapping sample sets. A total of 61 genome-wide (28 also study-wide) significant peaks observed in our analysis are located at loci harboring putative craniofacial genes (implicated from human malformations or animal models), but which had not yet been observed in GWAS for normal-range facial morphology. Our GWAS additionally revealed 53 genome-wide-significant (26 also study-wide-significant) peaks at loci harboring genes with no previously known role in facial development or disease. The annotation for each GWAS peak can be found in Supplementary Table 3.

**Genomic regions harboring multiple lead SNPs.** With our pheno- typing and analysis framework, in many cases we are able to provide a more nuanced understanding of the underlying genetic architecture of facial variation. For example, variants at the **TBX15-WARS2**
locus (1p12; Fig. 4) were previously reported to be associated with forehead prominence and self-reported chin dimples, indicating that this locus has multiple spatially separated effects on the face. In our current analysis, we see the same influence on forehead morphology, as previously reported by our group, with lead SNP rs3936018, located in the promoter region of WARS2, reaching its lowest significance in segment 14 ($P_{\text{MetaUK}}$(segment 14) = 8.01×10^{-49}). Interestingly, this lead SNP overlaps in location with a SNP not originally identified in our peak selection approach, rs12027501 ($P_{\text{MetaUK}}$(segment 1) = 1.03×10^{-43}). The latter was most significant in segment 1, the full face, and is not a good proxy for the former ($r^2$, 0.075; normalized coefficient of linkage disequilibrium $D'$, 0.979), indicating it is likely an independent statistical signal. Another signal, approximately 275 kb upstream of TBX15 (rs7513680), was most significantly associated with morphology in segment 51 ($P_{\text{MetaUK}}$(segment 51) = 7.03×10^{-13}), representing the cheek area around the corners of the mouth. Lastly, another GWAS peak is present approximately 301 kb downstream of WARS2 (rs17023457) with an effect in the upper cheeks ($P_{\text{MetaUK}}$(segment 48) = 3.26×10^{-13}). Of interest, we observed 24 such loci with multiple genome-wide-significant peaks that are each associated with different facial traits (Supplementary Table 5 and Supplementary Data 1).

**Genetic interactions impacting facial variation.** To better analyze and rank the effects of multiple genotypes on a facial trait, we utilized structural equation modeling (SEM) to refine our understanding of which groups of genome-wide-significant variants best explain the variance observed in each facial segment. SEM is a multivariate statistical analysis technique that analyzes structural relationships between measured variables (for example, genetic variants and covariates) and latent constructs (univariate phenotypes derived from the PCs of the analyzed facial segment). This was done in an iterative manner, resulting in 50 well-fitting SEM models (corresponding to 50 facial segments; Supplementary Data 2). For each of these 50 models, the output included a univariate latent variable and a list of variants ranked by their estimated contribution to that variable, highlighting the polygenic nature of facial variation captured by the latent variable. Higher correlation of cross-sample H3K27ac activity was found when comparing SNPs deemed significant by the same SEM model than when comparing SNPs nonsignificant in the same SEM model (Extended Data Fig. 7). Additionally, of the SEM-significant SNPs, four SNP combinations displayed evidence of pairwise epistatic interactions (Table 1, Fig. 5, Extended Data Fig. 8 and Supplementary Note 3).

**Discussion** In their review, Atchley and Hall provided a framework with which we can better understand and describe the development of complex morphological structures. In this analysis, we have focused on one part of this framework and have identified intrinsic genetic factors contributing to normal-range variation in the structure of the human face. By implementing an open-ended multivariate association method, in which the inherent morphological variation within each of these segments drives the association, and by using both standard and modified-for-multivariate follow-up bioinformatic approaches, we describe the association between SNPs and facial traits as well as the likely cellular functions of the regions.
surrounding these SNPs. We also highlight regions with multiple SNPs affecting different facial phenotypes as well as evidence for multiple SNPs working in concert to produce a single phenotype. Taken in summary, our results illustrate an avenue for investigating the coordinated processes underlying complex morphological structures, like the human face, at a deeper level than single associations between genotype and univariate phenotype.

Overall, our association results reflect patterns from known biological processes. For instance, linkage disequilibrium (LD) score regression correlations between segments seem to reflect the shared embryological origins of different parts of the face, indicating that the hierarchical spectral clustering of the face based on structural correlations effectively partitions underlying genetic signals into biologically coherent groups. It is additionally clear from the large number of genome-wide-significant SNPs reaching their strongest association in the full face and segment 2 (covering the nose and upper lip) that these facial regions are ‘hotspots’ for genomic signals (Fig. 1). In general, quadrant II (representing the nose) and quadrant IV (representing the forehead and eyes) had the most genome-wide-significant lead SNPs reaching lowest significance in segments within each quadrant. This is unsurprising, given the close relationship between visible facial features in those areas and

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**Fig. 4 | TBX15-WARS2 multi-peak locus.** LocusZoom plots and facial effects for four association signals near the TBX15-WARS2 locus. Clustering based on *r*² was performed to separate noncorrelated signals, resulting in the separation of four SNP clusters. Color for each SNP is based on cluster association, with saturation indicating *r*² correlation with the most significant SNP in the cluster. SNPs represented by diamonds are the genome-wide-significant lead SNPs also present in the 1000G Phase 3 dataset; SNPs represented by circles are adjacent SNPs also present in the 1000G Phase 3 dataset; SNPs represented by asterisks are those not present in the 1000G Phase 3 dataset. For the segment in which each lead SNP had its lowest effect, we plot the facial effects for the lead SNPs reaching significance in that segment as the normal displacement (displacement in the direction normal to the facial surface) in each quasi-landmark going from minor to major allele, with red colored areas shifting outward while blue colored areas shift inward.
Indeed, regions with less correspondence to underlying skeletal structure, like the upper lip (quadrant I), had many fewer lead SNPs reaching lowest significance in the contained segments, and facial regions with some structural correspondence but still greatly impacted by age and adiposity, like the lower face and cheeks (quadrant III), had only slightly more.

Reassuringly, the genes located within 500 kb of our genome-wide-significant lead SNPs were highly enriched for processes and phenotypes associated with craniofacial development and morphogenesis in humans and mice (Extended Data Fig. 5).

Notably, the top human phenotype was oral clefting, indicating a substantial overlap between the genes involved in normal facial variation and those implicated in the most common craniofacial birth defect in humans. Furthermore, many of the surrounding genes to which the genome-wide-significant lead SNPs were annotated are known to be involved in pathways relevant for craniofacial development, such as the Wnt signaling and TGFβ pathways (Extended Data Fig. 5b). Our GWAS signals were also enriched for processes associated with limb development and related phenotypes, pointing to a shared genetic architecture between faces and limbs (Extended Data Fig. 5a) and a number of genes near our genome-wide-significant loci (for example, Dlx homeobox genes, BMP genes, and FGFR2).

**Table 1 | Four SNPs with evidence of epistatic interactions**

| Segment | SNP 1 | SNP 2 | Test statistic | P value |
|---------|-------|-------|----------------|---------|
| rsID    | Location     | Gene annotation | rsID    | Location     | Gene annotation | statistic | P value |
| 6       | rs10838269   | 11:44378010   | ALX4   | rs11175967   | 12:66321344   | HMGA2   | 23.9422   | 9.94×10⁻⁷   |
| 9       | rs76244841   | 1:2775953    | PRDM16 | rs62443772   | 7:42131949   | GLI3    | 16.5745   | 4.68×10⁻⁶   |
| 11      | rs6740960    | 2:42181679   | PKDC1  | rs6795164    | 3:133885925  | SLCO2A1 | 16.3707   | 5.21×10⁻⁵   |
| 22      | rs7373685    | 3:128107020  | GATA2  | rs7843236    | 8:121980512  | SNTB1   | 15.7837   | 7.10×10⁻⁵   |

For each of the 50 segments with a refined SEM model, we used the latent variables and SNP lists to test for evidence of epistasis using a two-sided linear regression epistasis test in Plink v.1.9, with Bonferroni multiple-testing correction. For the four SNP pairs with significant evidence of epistatic interactions, Table 1 lists the epistasis P value, rsID, GRCh37 location and gene annotation. The phenotypic and marginal distributions for the pairs are depicted as boxplots in Fig. 5 and Extended Data Fig. 8.
have well-established roles in limb development. These findings are also supported by the large number of human syndromes that present with both facial and limb malformations.

For the regions surrounding the 203 genome-wide-significant lead SNPs, both CNCCs and embryonic craniofacial tissues showed the highest enrichment in H3K27ac signal (Fig. 2a). These observations are consistent with (1) activity of our 203 genome-wide-significant lead SNPs in CNCCs and embryonic craniofacial tissues and (2) an embryonic origin for human facial variation across the timeline of facial development, as CNCCs represent an early time point in facial development whereas the craniofacial tissues represent progressively later timepoints. In both CNCCs and craniofacial tissue at all sampled developmental stages, regions in the vicinity of the 203 genome-wide-significant lead SNPs were significantly enriched for predicted enhancers and not promoters. This is an especially intriguing result, as recent evidence has described the action of multiple enhancers, each showing different tissue or timing specificity, in modulating expression levels to affect craniofacial development. Complementing our GREAT analysis results, indicating that some genes near our GWAS peaks are involved in both facial and limb development, a subset of genome-wide-significant lead SNPs showed preferential activity in additional in-vitro-derived cell types relevant to both the face and the rest of the skeletal system, including osteoblasts, chondrocytes, differentiating skeletal muscle myoblasts, fibroblasts and keratinocytes (for example, cluster 3; Fig. 3). Together, these results suggest that genetic variation underlying facial morphology operates by modulating enhancer activity across multiple cell types throughout the timeline of embryonic facial development.

A total of 61 genome-wide-significant peaks from our analysis did not overlap with the results of prior GWAS for normal-range facial morphology, but were located nearby putative craniofacial genes implicated from human malformations or animal models. For instance, Msx1 has been implicated in orofacial clefting in humans and mice, and is also expressed widely in lip and dental tissues during development. We observed two distinct peaks at the Msx1 locus (4p16.2), one approximately 55 kb upstream of Msx1 with a pronounced effect on the lateral upper lip (lead SNP rs13117653; \(P\_\text{meta-l}4\)(segment 34) = 4.2 × 10\(^{-10}\)) and a second peak, about 323 kb upstream of Msx1 and located in the intron of STX18, involving the lower lateral lip and mandible (lead SNP rs3910659; \(P\_\text{meta-l}4\)(segment 25) = 4.45 × 10\(^{-8}\). Extended Data Fig. 9a–e). This result could indicate a potential role of STX18 in craniofacial development, although the STX18 protein is important primarily for functioning of the endoplasmic reticulum. Alternatively, this result could provide further evidence that complex phenotypic effects seen in our human sample could be due to the action of multiple regulatory elements within a single locus. In support of this, Attanasio et al. demonstrated that the activity of Msx1 in the second pharyngeal arch and maxillary process of the e11.5 mouse embryo is recapitulated by the combined activity of two separate enhancers.

We also identified 53 genome-wide-significant signals in regions harboring genes with no previously known role in craniofacial development or disease, although many of the implicated genes are known to have a general role in developmental processes critical to morphogenesis. For example, in the current study, variants at the DACT1 locus are associated with mandibular morphology (Extended Data Fig. 9f–h). DACT1 is an established antagonist of the Wnt signaling pathway, which is known to be involved in craniofacial development, although DACT1 is studied mostly for its involvement in gastric cancer. However, DACT1 has also been shown to inhibit the delamination of neural crest cells, further supporting its involvement in facial development. These new signals are promising new candidates for potential roles in facial morphogenesis.

In addition to better understanding which parts of the face had the most signals, we capitalized on the utility of facial segmentation via hierarchical clustering to finely parse out the effect of a SNP even within a complex genomic region. Notably, we observed 24 loci with multiple genome-wide-significant peaks each associated with different facial traits, suggesting that these variants might overlap with or be impacted by regulatory elements that affect the face in highly specific ways (Supplementary Table 5 and Supplementary Data 1). An important consideration to our peak selection procedure is that it is statistical and heuristic in nature, being based on investigator-chosen thresholds of both distance and similarity of associated facial phenotypes, and thus is not perfect. Refining a peak selection approach based on combinations of distance, LD patterns, and trait similarity was beyond the grasp of this paper, but we believe such an approach has potential for further interrogating the complex genetic architecture of facial variation, as we have illustrated using the TBX15-WARS2 locus (Fig. 4).

Given the complexity of the human face and its component traits, it is likely that the genetic architecture contributing to facial variation includes groups of genomic regions that contribute to the same facial trait, perhaps through actions in similar cell types or explicit interactions among variants. Importantly, genome-wide-significant SNPs that significantly explained variance in the same segment, based on the SEM for that segment, showed higher correlations of cross-sample H3K27ac activity than when compared with SNPs that did not, indicating that the SEM-refined lists of SNPs for each segment are likely those that are similar in either their spatial or temporal cellular activity (Extended Data Fig. 7). Tests for epistasis using the SEM-refined SNP lists for each segment identified four SNP combinations with significant evidence of pairwise epistatic interactions (Table 1). For example, rs76244841 (PRDM16 associated; \(P\_\text{meta-l}4\)(segment 30) = 1.48 × 10\(^{-9}\)) and rs62443772 (GLI3 associated; \(P\_\text{meta-l}4\)(segment 22) = 5.35 × 10\(^{-16}\)) were found to have a significant interaction in facial segment 9, which covers the premaxillary soft tissue from the base of the columella to the oral commissure (Table 1 and Fig. 5). Interestingly, PRDM16 and GLI3 are both part of a tetrameric Hedgehog signaling complex in Drosophila melanogaster (Supplementary Note 3\(^{16,29}\)). Overall, these results indicate that the statistical evidence of SNP groups influencing polygenic facial variation identified through SEM, and explicit variant interactions suggested by the epistasis analysis, are potentially representative of true biological relationships but must be confirmed with further study.

In conclusion, with this work we have not only reported genomic variants influencing normal-range facial variation, but have also sought to use our in-depth facial phenotyping approach and bioinformatic tools to illustrate one way in which researchers without access to functional follow-up analyses can delve deeper into the genetic architecture of complex morphological traits. These results illustrate the potential to highlight spatial and temporal connections between SNPs, representing a major step forward in our ability to characterize the polygenic genetic architecture of complex morphological structures. In performing an open-ended and minimally restrictive study, we are optimistic that our results will be useful for other research efforts to better understand the biological forces that shape human and nonhuman morphology.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at [https://doi.org/10.1038/s41588-020-00741-7](https://doi.org/10.1038/s41588-020-00741-7).

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Methods
Sample and recruitment. The samples used for analysis included a combination of three independently collected datasets from the United States (US; \( n = 4,680 \)) and one dataset from the United Kingdom (UK; \( n = 3,566 \)), for a total sample size of \( n = 8,246 \). The US samples originated from the 3D Face Norns cohort (JDFN) and studies at the Pennsylvania State University (PSU) and Indiana University-Purdue University Indianapolis (IUPUI). The UK dataset included samples from the Avon Longitudinal Study of Parents and their Children (ALSPAC)\(^{55-57} \). Institutional review board approval was obtained at each recruitment site, and all participants gave their written informed consent before participation. For children, written consent was obtained from a parent or legal guardian. Some individuals from the 3DFN and PSU samples were tested previously for associations with facial morphology in our prior work\(^{58} \). A breakdown of the samples used for analysis is shown in Supplementary Table 2 and further details are available in the Supplementary Methods. In all datasets, participants with missing information in sex, age, height or weight, or with insufficient image quality were removed.

Genotyping and imputation. Due to the several genotyping platforms used for the US cohort (details in the Supplementary Methods), we chose to impute the samples from each platform separately, then combine the imputed results\(^{59} \). For each dataset, standard data cleaning and quality assurance practices were performed based on the GRCh37 genome assembly. Phasing was performed using SHAPEIT2 (v.2r900)\(^{60} \) and imputation to the 1000G Phase 3 reference panel\(^{61} \) performed using the positional Burrows-Wheeler Transform pipeline (v.3.1) of the Sanger Imputation Server (v.0.0.6)\(^{62} \). After post-imputation quality control and intersection of imputed SNPs, a single merged dataset of all US participants was created with 7,417,619 SNPs for analysis.

The raw genotype data from ALSPAC were not available, and restrictions are in place against merging the ALSPAC genotypes with any others. For this reason, ALSPAC genotypes, phased using SHAPEIT2\(^{63} \) and imputed to the 1000G Phase 1 reference panel (Version 3\(^{64} \) using IMPUTE2\(^{65} \), were obtained directly from the ALSPAC database and held separately during the analysis. After post-imputation quality control, the ALSPAC dataset contained 8,629,873 SNPs for analysis.

For both datasets, SNPs on the X chromosome were coded 0/2 for hemizygous males, to match with the 0/1/2 coding for females\(^{66} \).

Ancestry axes and selection of European participants. From the post-imputation merged dataset of US participants, we identified the European participants by projecting them into a PC space constructed using the 1000G Phase 3 dataset, first filtered for LD and SNPs shared between both datasets. Further details are available in the Supplementary Methods. In the combined PC space, we calculated the ancestry axes for the US participants and the Euclidean distance between all US participants and the 1000G samples. Using a 4th nearest neighbor algorithm, we identified the five nearest 1000G neighbors for each US participant. The most common 1000G population label from these five nearest neighbors was then assigned to the US participant, and participants assigned the 1000G European population labels were selected for analysis.

Ancestry axes were calculated for the UK participants by projecting them into the 1000G Phase 3 dataset in a manner similar to that described for the US participants. Since all ALSPAC participants available for this analysis were European, no additional ancestry refinement was performed.

3D image acquisition. For all datasets, 3D images were captured using either a digital facial stereophotogrammetry system or a laser scanning system. All participants were asked to have closed mouths and to maintain a neutral facial expression during image capture\(^{67} \). For the 3DFN sample, facial surfaces were acquired using the 3DMDface (3DMD) camera system. PSU images were obtained with either the 3dMDface or Vectra H1 system (Canfield Scientific). The IUPUI sample was fully imaged using the Vectra H1. The ALSPAC sample was imaged using a Konica Minolta Vivid 900 laser scanner (Konica Minolta Sensing Europe). For this system, two high-resolution facial scans were taken and then processed, merged and registered using a macro algorithm in Rapidform 2004 software (INUS Technology Inc.).

3D image registration and quality control. The 3D surface images and their reflections were registered using the MeshMonk registration framework (v.0.0.6)\(^{68} \) in Matlab 2017b. This process results in a homologous configuration of 7,160 males, to match with the 0/1/2 coding for females\(^{12} \). This process results in a homologous configuration of 7,160 males, to match with the 0/1/2 coding for females\(^{12} \). The strength of covariation between quasi-landmarks was defined using Escoufier’s RV coefficient\(^{69} \). The RV coefficient was then used to build a structural similarity matrix that defined the hierarchical construction of 63 facial segments, broken into five levels (Extended Data Fig. 1a). The configurations of each segment were then subjected independently to a Generalized Procrustes analysis\(^{70} \), after which a PCA was performed in combination with parallel analysis to capture the major variance in the facial segments with fewer variables\(^{71-73} \) (Extended Data Fig. 1b).

Multivariate genome-wide association meta-analyses. The meta-analysis framework utilized consists of three steps performed separately for each of the 63 segments: identification, verification, and meta-analysis (Extended Data Fig. 2). For all analyses, the genotypes were coded additively based on the presence of the major allele. In the identification step, for each of the 63 facial segments, each SNP was associated with phenotypic variation using CCA (canoncorr in Matlab 2017b). CCA is a multivariate analysis that extracts the linear combination of PCs, which represent the direction of phenotypic effect in shape space (which we call a ‘trait’) that are maximally correlated with a SNP and returns a correlation value between those PCs and the SNP tested. Because CCA does not accommodate adjustments for covariates, we removed the effect of relevant covariates (sex, age, age-squared, height, weight, facial size, the four genome ancestry axes and 206 PCA tracks). For this system, two high-resolution facial scans were taken and then processed, merged and registered using a macro algorithm in Rapidform 2004 software (INUS Technology Inc.).

For this system, two high-resolution facial scans were taken and then processed, merged and registered using a macro algorithm in Rapidform 2004 software (INUS Technology Inc.).

CCA-US and CCA-UK meta-analyses were separately adjusted for sex, age, age-squared, height, weight, facial size, the four genome ancestry axes and 206 PCA tracks. Per SNP, the lowest \( P \) value was obtained with the Student’s \( t \) distribution function (function tcdf in Matlab 2017b).

In the meta-analysis step, the identification step identifies the phenotypic trait most correlated with each SNP (\( P \) (CCA-US) and \( P \) (CCA-UK)) representing the strength of the correlation. CCA has also been implemented in ‘mv-PLINK’\(^{74} \). Performance tests of mv-PLINK have shown that it outperforms univariate methods and has similar power to other multivariate methods of association\(^{75-77} \), which generally have higher statistical power than univariate methods\(^{78-80} \).

In the meta-analysis step, the identification \( P \) value (from Rao’s \( F \)-test on the canonical correlation) and the verification \( P \) value (from the univariate regression) were combined using Stouffer’s method\(^{28,29} \), which was chosen because a meta-analysis of beta values was not possible given that the CCA returns a positive correlation value, not a beta statistic. The entire process was repeated, resulting in two meta-analysis \( P \) values (\( P \)CCA-US and \( P \)CCA-UK) for each of the two identified traits per segment and per SNP. First, using US data in the identification stage and UK data as verification (META-US or US-driven), then using UK data in the identification stage and US data as verification (META-UK or UK-driven). A validation of our analysis pipeline is available in Supplementary Note 1.

Sharing of genome-wide signal between facial segments. To assess the extent to which genome-wide signals of association with facial variation were shared between a pair of facial segments, LD score regression\(^{81,82} \) was applied to the meta-analysis, after converting the meta \( P \) values to \( z \)-scores and ignoring the sign or direction of effect. The former was required because of the multivariate nature of our results and the latter was needed since CCA is a one-sided test with canonical correlations always between \([-1,1] \). As a result, all resulting genetic correlations reported here are restricted to be positive as well. Further details on the calculation of LDSC values are available in the Supplementary Methods. This process was done twice, once each for the US- and UK-driven meta-analyses. A high degree of congruence (\( r = 0.95 \)) between the results based on the US- and UK-driven meta-analyses was observed, and the average correlation of both between each pair of facial segments was reported. The 63 x 63 matrix of average correlations was visualized on top of the facial segmentation hierarchy to assess correlation both within and between facial quadrants (Extended Data Fig. 3) and used to perform average-linkage hierarchical clustering (Extended Data Fig. 4).

GWAS peak selection. The analysis strategy yielded 126 meta-analysis \( P \) values and 126 traits for every SNP, representing the 63 segments x two meta-analysis tracks. Per SNP, the lowest \( P \) value was selected, and we noted in which meta-analysis track (META-US or META-UK) and which meta-analysis track (‘Best meta-analysis track’) and segment (‘Best segment’) this \( P \) value occurred. The study-wide Bonferroni threshold size, the first four genomic ancestry axes and the camera system, using PLSR (function plscore from Matlab 2017b). As an illustration, the age adjustment is visualized in Supplementary Fig. 2. After adjustment, facial segments were defined by grouping vertices that are correlated strongly using hierarchical spectral clustering\(^{83,84} \). The strength of covariation between quasi-landmarks was defined using Escoufier’s RV coefficient\(^{4,4} \). The RV coefficient was then used to build a structural similarity matrix that defined the hierarchical construction of 63 facial segments, broken into five levels (Extended Data Fig. 1a). The configurations of each segment were then subjected independently to a Generalized Procrustes analysis\(^{4} \), after which a PCA was performed in combination with parallel analysis to capture the major variance in the facial segments with fewer variables\(^{71-73} \) (Extended Data Fig. 1b).
Lists of human CNCC regulatory elements were annotated based on multiple by silhouette width (Fig. 3). For the initial peak selection, we chose to group SNPs below genome-wide threshold by genomic position, and the SNP with the lowest P value per genomic region was selected as the lead SNP. Within a ±500-kb window of the resulting genome-wide-significant lead SNPs, we further refined the selection by performing a regression on the predefined and refined identification stage (in Best meta-analysis track and Best segment) to determine if adjacent SNPs showed consistent effects with the lead SNP resulting in 218 genome-wide-significant lead SNPs. Of these 218 lead SNPs, 203 showed consistent traits in the US and UK datasets in the Best segment (Supplementary Table 3), with more details available in Supplementary Note 2 and the permutation outcomes available in the FigShare repository for this article. Although a study-wide threshold was calculated, we chose to annotate lead SNPs reaching at least genome-wide threshold to retain as many potently biologically meaningful results as possible. The FigShare repository also provides information on all SNPs reaching suggestive significance ($P \leq 6.96 \times 10^{-4}$) was calculated as $5 \times 10^{-15}$ ([1.0042 $\times 1.6631 \times 4.1345$], with the denominator values representing the number of independent tests per SNP across both meta-analysis tracks, and across all segments, respectively. These values were calculated using the R package herdm (v.23.0.1), which reports a two-sided $P$ value. For our analyses, separate SEM models were constructed for each segment using each of the 203 genome-wide-significant lead SNPs and the shape PCs for all participants, with additional information available in the Supplementary Methods.

For each of the 50 SEM models where the refinement process was successful (details in the Supplementary Methods), final model fit indices and model parameter estimates are provided in Supplementary Data 2. Reassuringly, for segments that are closely related in the segmentation hierarchy (segments 5, 11, 23 and 47) there is an average overlap of 46% of the variants meeting the $P < 0.05$ cutoff for SEM significance, compared to 13.6% average overlap for nonhierarchically related segments (segments 5 and 6). The H3K27ac activity across all cell types was compared for significant variants both within and between segments using Spearman’s rho using two-sided Kruskal–Wallis tests (Extended Data Fig. 7).

Epistasis analysis. We additionally used the univariate latent variable and the variants passing the $P < 0.05$ significance cutoff from the final refined SEM models (P < 0.1 for segments 7, 16 and 25) to assess whether interactions between genotype increase or decrease the distribution of the latent variable. For each segment, the effect on the latent variable of all diplotype combinations of variants was assessed via a linear regression epistasis analysis Plot in Plink v.1.9 (ref. 84). At a correlation threshold for epistats of r = 0 (P < 0.05 Table 1). For these four pairs, the nine diplotype combinations and their normalized posterior and marginal distributions were plotted (Fig. 5 and Extended Data Fig. 8) to assess the genotypic contribution to epistatic masking (the combination of two variants reduce the phenotype) and boosting (the combination of two variants increase the phenotype). For each diplotype combination, the marginal SNP’s genotypic medians were averaged to visualize the predicted phenotypic distribution that would occur if the two genotypes were acting independently, and this average median was compared to the medians of the combined diplotypes. Significance testing was performed using a two-sided Mood’s Median test with one degree of freedom. These steps were performed using the R packages agricolae (v.1.3-0), covtest (v.1.0.8), gplplot (v.3.1.1), ggsplot2 (v.2.2.1), gridExtra (v.2.3.6), grid (v.3.6.2), htmlTable (v.4.2.0), psych (v.1.8.12) and data.table (v.1.12.0).

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. All of the genotypic markers for the 3DFN dataset are available to the research community through the dbGaP controlled-access repository (http://www.ncbi.nlm.nih.gov/gap) at accession no. phs000949.v1.p1. The raw source data for the phenotypes—the 3D facial surface models in.obj format—are available through the FaceBase Consortium (https://www.facebase.org) at accession no. F008009/491/01. Access to these 3DF facial surface models requires proper institutional ethics approval and approval from the FaceBase data access committee. Additional details can be requested from S.W.

The participants making up the PSU and IUPUI datasets were not collected with broad data sharing consent. Given the highly identifiable nature of both facial images and associated information and unresolved issues regarding risk to participants, we opted for a more conservative approach to participant recruitment. Broad data sharing of the raw data from these collections would thus be in legal and ethical violation of the informed consent obtained from the participants. This restriction is not because of any personal or commercial interests. Additional details can be requested from M.D.S. and S.W. for the PSU and IUPUI datasets, respectively. The ALSAPAC (UK) data will be made available to bona fide researchers on application to the ALSAPAC Executive Committee (http://www.bris.ac.uk/alspac/researchers/data-access). Ethical approval for the study was obtained from the ALSAPAC Ethics and Law Committee and the Local Research Ethics Committees. Publicly available data used were the 1000G Phase 3 data (ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/), the list of HapMap 3 SNPs excluding the MHC region (http://ldsc.broadinstitute.org/static/media/w_hm3_noMHC.snplist.zip), and Chi2–pseq files from Prescott et al. (GSE70575), Najafzadeh et al. (GSE82295), Baumgart et al. (GSE81979), Nott et al. (https://genome.ucsc.edu/;v/nottalex/glass_Lab_BrainCellTypes_hg19), Prescott et al. (GSE119977), Garcia-Compean et al. (GSE41795), and the Epigenomics Project (https://egg2.wustl.edu/roadmap/data/byFileType alignments/consolidated/). Meta-analysis GWAS statistics are available on GWAS Catalog (GCP000044). All data relevant to run future replications and meta-analysis efforts are provided in the FigShare repository for this work, along with additional figures (https://doi.org/10.6084/m9.figshare.c.6467216). Items 4 and 5 of the Supplementary Methods apply to the anthropometric mask used; (2) association statistics and effects of the 203 lead SNPs, facia levels, LocusZoom plots and association statistics from each stage of the analysis for the 203 lead SNPs; (3) calculation of study-wide significance.
threshold: script and permutation outcomes needed to replicate the calculation of the study-wide-significance threshold; (4) facial segment assignments: segment assignments for each quasi-landmark in the anthropometric mask; (5) Fig. 2a: labeled a version of Fig. 2a, with all cell types and tissues labeled; (6) GREAT Export: raw output of the GREAT analysis; (7) PCA shape constructs: PCA shape spaces for all 63 facial segments; (8) QQ plots: QQ plots for each segment in all stages of the analysis; (9) script to explore facial segments and GWAS hits: MatLab script for select data exploration functions; (10) SNPs reaching suggestive significance in either analysis track: association statistics of all SNPs with P ≤ 5 x 10−5 in METAUS or METAUK tracks; (11) source data for manuscript figures: source data in Excel format for all figures, where possible.

Code availability

KU Leuven provides the MeshMonk (v.0.0.6) spatially dense facial-mapping software, free to use for academic purposes (https://github.com/TheWebMonks/meshmonk). Matlab 2017b implementations of the hierarchical spectral clustering to obtain facial segmentations are available from a previous publication (https://doi.org/10.6084/m9.figshare.7649024).

The statistical analyses in this work were based on functions of the statistical toolbox in Matlab 2017b, SHAPETZ (v.2.1900), Sanger Imputation Server (v.0.0.6), PBWT pipeline (v.3.1), MeshMonk (v.0.0.6), LDSC (v.1.0.1), FUMA (v.1.3.3), GREAT (v.1.0.9), lajanav (v.0.6–6.3), R (v.3.4), agricolae (v.1.3-0), cowplot (v.1.0.0), ggplot2 (v.3.1.1), ggpubr (v.0.2), gridExtra (v.2.3.2), ggplot (v.3.0.3), grid (v.3.6.2), Hmisc (v.4.2–0), data.table (v.1.12.0), Genotype Harmonizer (v.1.4.20), KING (v.2.1.3), bowtie2 (v.2.3.4.2), bedtools (v.2.27.1) and Bioconductor (v.3.7), as mentioned throughout the References.

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Author contributions

P.C., M.D.S., S.M.W., J.R.S., J.W. and S.W. conceptualized the study (ideas; formulation or evolution of overarching research goals and aims), J.D.W., K.I., R.E., M.K.L., J.L., S.W. and P.C. carried out the data curation (management activities to annotate (produce related metadata and maintain research data for initial use and later re-use), J.D.W., K.I., S.N., R.E., H.H., J.R., J.L. and P.C. carried out the formal analysis (application of statistical, mathematical, computational or other formal techniques to analyze or
synthesize study data). S.R., H.L.N., E.E., T.S., M.L.M., J.R.S., J.W., S.W., S.M.W., M.D.S. and P.C. were responsible for funding acquisition (acquisition of the financial support for the project leading to this publication). J.D.W., K.I., S.N., R.J.E., H.H., J.R., M.K.L., J.L. and P.C. carried out the investigation (conducting a research and investigation process, specifically performing the experiments or data/evidence collection). J.D.W., S.N., R.J.E., J.M., S.R., E.E.Q., H.L.N., T.S., M.L.M., J.W., S.W., S.M.W. and M.D.S. provided the resources (provision of study materials, computing resources or other analysis tools). P.C., S.M.W., M.D.S., S.W., J.W., J.R.S., M.L.M., T.S., H.P. and G.H. carried out the supervision (oversight and leadership responsibility for the research activity planning and execution, including mentorship external to the core team). J.D.W., K.I., S.N., R.J.E., H.H., J.R., M.K.L. and P.C. did the visualization (preparation, creation and/or presentation of the published work, specifically visualization/data presentation). J.D.W., K.I., S.N., R.J.E. and J.R. wrote the original draft. J.D.W., K.I., S.N., R.J.E., H.H., J.R., S.R., E.E.Q., M.L.M., H.P., J.R.S., J.W., S.W., S.M.W., M.D.S. and P.C. reviewed and edited the final manuscript.

Competing interests
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Additional information
Extended data is available for this paper at https://doi.org/10.1038/s41588-020-00741-7.
Supplementary information is available for this paper at https://doi.org/10.1038/s41588-020-00741-7.
Correspondence and requests for materials should be addressed to J.D.W., K.I. or P.C.
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Extended Data Fig. 1 | Hierarchical spectral clustering of facial shape. a, Global-to-local facial segmentation of all 3D images (nTotal = 8,246), obtained using hierarchical spectral clustering. Segments are colored in teal and identical to those in Fig. 1. Roman numerals represent ‘quadrants’ of facial segments. b, The number of principal components retained after parallel analysis for each facial segment.
**Extended Data Fig. 2 | Study design.** Sample Wrangling: Images and genotypes from each study were intersected and unrelated participants of European ancestry, with quality-controlled images, covariates, and imputed genetic data were selected to obtain the analyzed data. Identification: For each facial segment, canonical correlation analysis (CCA) and Rao’s F-test approximation was used to identify the multivariate combination of facial principal components most correlated with the genotypes, which led to a P-value (PCCA-US or PCCA-UK) and multivariate phenotypic trait most correlated with each SNP (TraitUS and TraitUK). Verification: The principal components of the other dataset were then projected onto this trait to obtain a univariate variable representing the distribution of participants from the verification dataset for the trait identified in the identification dataset (UniVarUK and UniVarUS). The genotypes of the verification dataset are then tested against this variable via linear regression, resulting in an additional P-value (PUniVar-UK and PUniVar-US). Meta-Analysis: The P-values from identification and verification are meta-analyzed using Stouffer’s method, resulting in the final set of P values from each meta-analysis track (PMETA-US and PMETA-UK).
Extended Data Fig. 3 | Genomic signal correlations. LDSC correlations between segments. **a**, Correlations between segments from different quadrants, ranging from 0.8 to 0.88, which seem to reflect both physical proximity of segments on the face and shared embryological origins. **b**, Correlations ranging from 0.88 to 1, which are mostly between segments within the same facial quadrant.
Extended Data Fig. 4 | Clustering of facial segments on the basis of shared genetic signals. Correlations between facial segments on the basis of SNP $P$ values were calculated using LDSC, as described in Methods, and average-linkage hierarchical clustering was performed using the matrix of correlation values. Quadrant colors in legend refer to the quadrant of the polar dendrogram in which the facial segment lies in, also represented by the facial images at the top, and embryonic facial prominences are assigned to each facial segment.
Extended Data Fig. 5 | GREAT and FUMA analyses showing enrichment for craniofacial and limb development. a, GREAT analysis. For the top ten GO terms in each category, plotted is the binomial test Bonferroni-corrected $P$ value (red, negative values) and binomial region fold enrichment (blue, positive values). Behind every GO term, in parentheses we indicate the number of genes in the test set with the annotation (Observed) and the total number of genes in the genome with the annotation (Total), with the format (Observed/Total). Dashed line represents significance at $P = \log_{10}(0.05) = -1.3$. b, FUMA analysis, indicating the KEGG pathways that were significantly enriched in our results. Multiple pathways are relevant for craniofacial development. The right panel shows the genes that are involved in the pathways.
Extended Data Fig. 6 | H3K27ac signal is significantly different in 203 lead vs. 203 random SNPs for relevant facial tissues. For all cell types and tissues, each represented by a point above, the median difference between H3K27ac RPM signal between the 203 lead SNPs vs. 203 random SNPs was tested for significance using a two-sided Wilcoxon rank-sum test. The thin dashed line represents the 5% false discovery rate P value of 0.0094, using the Benjamini–Hochberg method. Relative to the random, MAF-matched SNPs, the lead SNPs are significantly enriched for H3K27ac signal in many cell types, with the highest magnitude differences being from CNCCs (blue) and embryonic craniofacial tissues (orange). Test statistics used to create this plot are available in Supplementary Table 4.
Extended Data Fig. 7 | Correlation of H3K27ac activity among SEM models. a, For all segments (aka ‘masks’), we compared the H3K27ac activity for significant SNPs from the refined SEM model for variation in that facial segment. Plotted is the Spearman’s rho correlation between pairs of SNPs significant in the same SEM model (‘Within Mask’); pairs of SNPs where one is from the SEM model and the other is not (‘Within To Out’), and where both SNPs in the pair are from a different SEM model (‘Out To Out’). Segments where the distribution of correlation across all cell types was significantly different (Benjamini–Hochberg adjusted $P < 0.05$) based on a two-sided Kruskal–Wallis test are indicated in black. b, For all cell types, the median correlation across all segments is plotted for each of the three SNP groupings. Significance between the means was determined using a two-sided Kruskal–Wallis test. Boxplots plot the first and third quartiles, with a dark black line representing the median. Whiskers extend to the largest and smallest values no further than 1.5 x the inter-quartile range from the first and third quartiles, respectively.
Extended Data Fig. 8 | Phenotypic and marginal distributions for diplotype combinations. For a random SNP pairing (a) and each significant epistasis pair (b–d), boxplots are plotted to visualize the epistatic effect on the phenotype. The marginal phenotypic medians of the singular genotypes (non-shaded boxplots) were used to calculate and visualize the predicted diplotype phenotypic distribution that would occur if the two genotypes were acting alone. The median phenotype was also calculated for each diplotype as the average of the marginal medians of the singular genotypes (blue dashed lines on the colored plots). This median was compared to the observed medians of the diplotypes (solid black lines; colored boxplots) via Mood’s Median test with one degree of freedom. Log-transformed $P$ values were used to color boxplots if there was a significant ($P < 0.05; \log(P) > 1.30$) difference between the expected phenotype of the combined genotype and observed diplotype. Boxplots plot the first and third quartiles, with a dark black line representing the median. Whiskers extend to the largest and smallest values no further than $1.5 \times$ the inter-quartile range from the first and third quartiles, respectively.
Extended Data Fig. 9 | MSX1 and DACT1 loci. LocusZoom plots for the two association signals nearby MSX1 (a), which has previously been implicated in orofacial clefting in humans and mice, and DACT1 (f), which is a novel result. Points represent one-sided $-\log_{10}(P)$ of the METAUK meta-analysis track for the facial segment illustrated in the normal displacement figures (b, d, g) and are colored based on linkage disequilibrium with the labeled SNP. Asterisks indicate genotyped SNPs and circles indicate imputed SNPs. Facial effects for the two association signals nearby MSX1: rs3910659 (b) and rs13117653 (d) and the signal nearby DACT1: rs10047930 (g). Effects are the normal displacement (displacement in the direction locally normal to the facial surface) in each quasi landmark of the lowest facial segment reaching genome-wide significance in METAUK going from the minor to the major allele. Blue indicates inward depression; red indicates outward protrusion. Yellow rosette plots depict the $-\log_{10}(P)$ of the meta-analysis $P$ value (one-sided, right-tailed) per facial segment in METAUK track. Black-encircled facial segments have reached genome-wide significance ($P = 5 \times 10^{-8}$). (c) rs3910659; (e) rs13117653; (h) rs10047930.
Extended Data Fig. 10 | Regions nearby previously published SNPs associated with risk for Crohn’s disease are preferentially active in immune cells and tissues. Each boxplot represents the distribution of H3K27ac signal in 20 kb regions around 619 Crohn’s disease-associated SNPs from the NCBI-EBI GWAS catalog in one sample. See Methods for details on calculation of H3K27ac signal. Samples corresponding to immune cells and tissues are highlighted in red. Thin dashed line at -2.9 is the median level of signal across all cell types and tissues. Boxplots plot the first and third quartiles, with a dark black line representing the median. Whiskers extend to the largest and smallest values no further than 1.5 × the inter-quartile range from the first and third quartiles, respectively.
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Give P values as exact values whenever suitable.

For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings

For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes

Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

Three-dimensional images composed of surface and texture maps were taken using the 3dMD Face (3dMD, Atlanta, GA) and Vectra H1 (Canfield Scientific, Parsippany, NJ) 3D imaging systems, or the Konica Minolta Vivid 900 (Konica Minolta Sensing Europe, Milton Keynes, UK) laser scanner. Images gathered using the Konica laser scanner were processed using a macro algorithm in Rapidform 2004 software (INUS Technology Inc., Seoul, South Korea). Genotyping was performed using the Illumina OmniExpress + Exome v1.2 array, the 23andMe v3 and v4 arrays (Mountain View, CA), the Illumina Infinium Multi-Ethnic Global-8 v1 array, or the Illumina Human Hap550 quad array.

Data analysis

KU Leuven provides the MeshMonk spatially dense facial mapping software (v0.0.6), free to use for academic purposes (https://github.com/TheWebMonks/meshmonk). Matlab 2017b implementations of the hierarchical spectral clustering to obtain facial segmentations are available from a previous publication (https://doi.org/10.6084/m9.figshare.7649024). The statistical analyses in this work were based on functions of the statistical toolbox in Matlab 2017b, SHAPEIT2 (v2.r900), Sanger Imputation Server (v0.0.6), PBWT pipeline (v3.1), MeshMonk (v0.0.6), LDSC (v1.2.1), GEM (v1.3.3), GREAT (v3.0.0), Plink 1.9, lavaan (v0.6-3), R (v3.4), agricolae (v1.3-0), cowplot (v1.0.0), ggplot2 (v3.1.1), ggpubr (v0.2.0), gridExtra (v2.2.1), grable (v0.2.0), grid (v3.6.2), Hmisc (v4.2-0), psych (v1.8.12), data.table (v1.12.0), Genotype Harmonizer (v1.4.20), KING (v2.1.3), bowtie2 (v2.3.4.2), bedtools (v2.27.1), and biocductor (v3.7) as mentioned throughout the Methods.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.
Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

All of the genotypic markers for the 3DFN dataset are available to the research community through the dbGap controlled-access repository (http://www.ncbi.nlm.nih.gov/gap) at accession #phs000929.v1.p1. The raw source data for the phenotypes - the 3D facial surface models in .obj format - are available through the FaceBase Consortium (https://www.facebase.org) at accession #FB800000491.01. Access to these 3D facial surface models requires proper institutional ethics approval and approval from the FaceBase data access committee. Additional details can be requested from SMW [smwst646@pitt.edu].

The participants making up the PSU and IUPUI datasets were not collected with broad data sharing consent. Given the highly identifiable nature of both facial and genomic information and unresolved issues regarding risk to participants, we opted for a more conservative approach to participant recruitment. Broad data sharing of the raw data from these collections would thus be in legal and ethical violation of the informed consent obtained from the participants. This restriction is not because of any personal or commercial interests. Additional details can be requested from MDS [mds17@psu.edu] and SW [walshsus@iupui.edu] for the PSU and IUPUI datasets, respectively.

The ALSPAC (UK) data will be made available to bona fide researchers on application to the ALSPAC Executive Committee (http://www.bris.ac.uk/alspac/researchers/data-access). Ethical approval for the study was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committees.

KU Leuven provides the MeshMonk (v0.0.6) spatially dense facial mapping software, free to use for academic purposes (https://github.com/TheWebMonks/meshmonk). Matlab 2017b implementations of the hierarchical spectral clustering to obtain facial segmentations are available from a previous publication (https://doi.org/10.6084/m9.figshare.7649024.v1).

Publicly available data used were: 1000G Phase 3 (ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/), the list of HapMap 3 SNPs excluding the MHC region provided by LDSC (http://ldsc.broadinstitute.org/static/media/w_hm3.n0MHC.snplist.zip), and ChiP-seq files from Prescott et al. (GSE70751), Najafova et al. (GSE82295), Baumgart et al. (GSE89179), Nott et al. (https://genome.ucsc.edu/s/nottalexi/glassLab_BrainCellTypes_hg19), Pattison et al. (GSE119997), Wilderman et al. (GSE97752) and the Roadmap Epigenomics Project (https://egg2.wustl.edu/roadmap/data/byFileType/alignments/consolidated/)

Meta-analysis GWAS statistics are available on GWAS Catalog (GCP000044). All relevant data to run future replications and meta-analysis efforts are provided in the FigShare repository for this work34, along with additional figures (https://doi.org/10.6084/m9.figshare.c.4667261). Items available in the FigShare repository are: (1) Anthropometric mask: a Matfile of the anthropometric mask used; (2) Association statistics and effects of the 203 lead SNPs: Facial effects, LocusZoom plots, and association statistics from each stage of the analysis for the 203 lead SNPs; (3) Calculation of study-wide significance threshold: Script and permutation outcomes needed to replicate the calculation of the study-wide significance threshold; (4) Facial segment assignments: Segment assignments for each quasi landmark in the anthropometric mask; (5) Figure 2A labeled: A larger version of Figure 2A, with all cell types and tissues labeled; (6) GREAT Export: Raw output of outcomes needed to replicate the calculation of the study-wide significance threshold; (4) PCA shape constructs: PCA shape spaces for all 63 facial segments; (8) QQ plots: QQ plots for each segment in all stages of the analysis; (9) Script to explore facial segments and GWAS hits: Matlab script for select data exploration functions; (10) SNPs reaching suggestive significance in either meta-analysis track: Association statistics of all SNPs with P < 5 × 10-7 in METAUS or METAUK tracks; (11) Source data for manuscript figures: Source data in Excel format for all figures, where possible.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size
The sample size was determined by the amount of 3DFN data available in the public facial data repository (Facebase.org), the amount of data available in the B2261 ALSPAC study, and by the number of individuals of European descent with genotype data and 3D facial images that were collected with informed consent as part of several studies based at The Pennsylvania State University and Indiana University Purdue University Indianapolis. More information is found in Methods.

Data exclusions
For both US and UK participants, outlier 3D facial images, likely caused by image mapping errors, were identified using two approaches. First, as described in prior work, outlier faces were identified by calculating z-scores from the Mahalanobis distance between the average face and each individual face. Faces with z-scores higher than two were manually investigated. Second, a score was calculated that reflects the missing data present in the image due to holes, spikes, and other mesh artifacts, which can be caused by facial hair or errors during the preprocessing steps. Images with high scores, indicating large gaps in the mesh, were manually investigated. During the manual check, the images were either classified as poor quality and removed or were preprocessed and mapped again.

For US cohorts, genotype samples were excluded if there was poor concordance of genetic and reported sex, evidence of chromosomal aberrations, missing genotype call rate > 10%, and heterozygosity values ±3 standard deviations from the sample mean.

For the 3DFN sample, 3D images and genotype data were obtained from the 3D Facial Norms repository. Recruitment was limited to
Individuals aged 3 to 40 years old and of self-reported European ancestry. Individuals were excluded if they reported a personal or family history of any birth defect or syndrome affecting the head or face, a personal history of any significant facial trauma or facial surgery, or any medical condition that might alter the structure of the face. The intersection of unrelated participants with quality-controlled images, covariates, and genotype data from individuals of European descent resulted in 1,906 individuals for analysis.

The PSU sample included 3D images and genotypes of participants recruited through several studies at the Pennsylvania State University. Individuals were excluded from the analysis if they were below 18 years of age and if they reported a personal history of significant trauma or facial surgery, or any medical condition that might alter the structure of the face. No restriction on ancestry or ethnicity was imposed during recruitment, but only individuals of European descent were used in this study. The intersection of unrelated European participants with quality-controlled images, covariates, and genotype data resulted in 1,990 individuals for analysis.

The IUPUI sample includes 3D images and genotypic data from individuals recruited in Indianapolis, IN and Twinsburg, OH. Individuals who were below 18 years of age were recruited if they had a parent or legal guardian’s signature. Similar to the PSU sample cohort, no restrictions were placed on the recruitment of participants, but only unrelated individuals of European descent, without significant facial injury or medical condition, and those meeting all quality control criteria were used in this study (n = 784).

The UK sample was derived from the ALSPAC dataset, a longitudinal birth cohort in which pregnant women residing in Avon with an expected delivery date between 1 April 1991 and 31 December 1992 were recruited. At the time, 14,541 pregnant women were recruited and DNA samples were collected for 11,343 children. Genome-wide data was available for 8,952 subjects and of the 82261 study, titled “Exploring distinctive facial features and their association with known candidate variants.” In addition to this, 4,731 3D images were available. UK genotype samples were excluded on the basis of genetic sex and reported gender mismatches, minimal or excessive heterozygosity, disproportionate levels of individual missingness (>3%), and insufficient sample replication (IBD <0.8). The intersection of unrelated participants of European ancestry with quality-controlled images, covariates, and genotype data included 3,566 individuals.

Blinding was not relevant to this study, as no treatment outcomes were assessed and data analysis procedures were standardized across all individuals. Two independent datasets were constructed from the sampling efforts of four different research centers, and the analysis was done by yet another research center.

Replication was achieved by proper separation of the data into identification and verification datasets, based on completely separate sampling, imaging, genotyping, and imputation. More information is found in Methods.

No randomization took place, group membership of identification and verification was determined by the separately obtained datasets available. Because canonical correlation analysis does not accommodate adjustments for covariates, we removed the effect of relevant covariates (sex, age, age-squared, height, weight, facial size, the first four genomic ancestry axes, and the camera system), on both the independent (SNP) and the dependent (facial shape pre segmentation) variables using partial least squares regression (plsgress from Matlab 2017b), and thus performed the canonical correlation analysis under a reduced model with residualized variables.

Reporting for specific materials, systems, and methods

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Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ✗   | Antibodies            |
| ✗   | Eukaryotic cell lines  |
| ✗   | Palaeontology         |
| ✗   | Animals and other organisms |
|      | Human research participants |
|      | Clinical data         |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ✗   | ChiP-seq              |
| ✗   | Flow cytometry        |
|      | MRI-based neuroimaging |

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)

Cranial neural crest cells (CNCCs) originated from WiCell (H9 ESC) and the Fred Gage laboratory (IPSC; Salk Institute), available from Prescott et al. (GSE70751)
Fetal osteoblast cell line, undifferentiated and differentiated, originated from Najafova et al. (GSE82295)
Mesenchymal stem cell-derived osteoblasts originated from Baumgart et al. (GSE89179)
Various brain cell types originated from Nott et al. (https://genome.ucsc.edu/s/nottalex/glassLab_BrainCellTypes_hg19)
Surface ectoderm samples originated from Pattison et al. (GSE119997)
Embryonic craniofacial tissue originated from Wilderman et al. (GSE97752)
All other fetal and adult cell tissues and cell types originated from the Roadmap Epigenomics Project (https://egg2.wustl.edu/roadmap/data/byFileType/alignments/consolidated/)

Sample type, ID, and URL for each cell type/tissue is available in Supplementary Table 4
**Human research participants**

Policy information about studies involving human research participants

**Population characteristics**

For the 3DFN sample, 3D images and genotype data were obtained from the 3D Facial Norms repository. The repository includes 3D facial surface images and self-reported demographic descriptors as well as basic anthropometric measurements from individuals recruited at four US sites: Pittsburgh, PA (PITT IRB PRO09060553 and R80405013); Seattle, WA (Seattle Children’s IRB 121107); Houston, TX (UT Health Committee for the Protection of Human Subjects HSC-DB-09-0508); and Iowa City, IA (University of Iowa Human Subjects Office IRB (200912764 and 200710721). Recruitment was limited to individuals aged 3 to 40 years old and of self-reported European ancestry. Individuals were excluded if they reported a personal or family history of any birth defect or syndrome affecting the head or face, a personal history of any significant facial trauma or facial surgery, or any medical condition that might alter the structure of the face. The intersection of unrelated participants with quality-controlled images, covariates, and genotype data resulted in 1,906 individuals for analysis (Female N = 1,172; Male N = 734). Average height of these participants was 163.43 cm (sd = 20.57 cm). Average weight of these participants was 64.33 kg (sd = 22.38 kg).

The PSU sample included 3D images and genotypes of participants recruited through several studies at the Pennsylvania State University and sampled at the following locations: Urbana-Champaign, IL (PSU IRB 131103); New York, NY (PSU IRB 45727); Cincinnati, OH (UC IRB 2015-3073); Twinsburg, OH (PSU IRB 2503); State College, PA (PSU IRB 44929 and 4320); Austin, TX (PSU IRB 44929); and San Antonio, TX (PSU IRB 1278). Participants self-reported information on age, ethnicity, ancestry, and body characteristics, and data were gathered on height and weight. Individuals were excluded from the analysis if they were below 18 years of age and if they reported a personal history of significant trauma or facial surgery, or any medical condition that might alter the structure of the face. No restriction on ancestry or ethnicity was imposed during recruitment, but only individuals of European descent were used in this study. The intersection of unrelated European participants with quality-controlled images, covariates, and genotype data resulted in 1,990 individuals for analysis (Female N = 1,380; Male N = 610). Age ranged from 18 to 88 years old. Average height of these participants was 168.75 cm (sd = 9.23 cm). Average weight of these participants was 73.88 kg (sd = 17.05 kg).

The IUPUI sample includes 3D images and genotypic data from individuals recruited in Indianapolis, IN and Twinsburg, OH (IUPUI IRB 1409306349). Participants self-reported information on age, height, weight, and ancestry at the time of the collection. Individuals who were below 18 years of age were included if they had a parent or legal guardian’s signature. Similar to the PSU sample cohort, no restrictions were placed on the recruitment of participants, but only n = 784 individuals of European descent and those meeting all quality control criteria were used in this study (Female N = 539; Male N = 245). Age ranged from 7 to 78 years old. Average height of these participants was 169.24 cm (sd = 11.30 cm). Average weight of these participants was 71.88 kg (sd = 18.65 kg).

The UK sample was derived from the ALSPAC dataset, a longitudinal birth cohort in which pregnant women residing in Avon with an expected delivery date between 1 April 1991 and 31 December 1992 were recruited. At the time, 14,541 pregnant women were recruited and DNA samples were collected for 11,343 children. Genome-wide data was available for 8,952 subjects of the B2261 study, titled “Exploring distinctive facial features and their association with known candidate variants.” In addition to this, 4,731 3D images were available along with information on sex, age, weight, height, ancestry, and other body characteristics. The ALSPAC study website contains details of all the data that is available through a fully searchable data dictionary (http://www.bris.ac.uk/alspac/researchers/our-data/). The intersection of participants of European ancestry with quality-controlled images, covariates, and genotype data included 3,566 individuals (Female N = 1,884; Male N = 1,682). Ethical approval for the study was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committees. Informed consent for the use of data collected via questionnaires and clinics was obtained from participants following the recommendations of the ALSPAC Ethics and Law Committee at the time. Consent for biological samples has been collected in accordance with the Human Tissue Act (2004). Age ranged from 14 to 17 years old. Average height of these participants was 169.38 cm (sd = 8.42 cm). Average weight of these participants was 61.52 kg (sd = 11.75 kg).

For all datasets, there was no imbalance in gender, and weight and height distributions follow typical distributions seen in a European-derived population. We removed the effect of relevant covariates (sex, age, age-squared, height, weight, facial size, the first four genomic ancestry axes, and the camera system), on both the independent (SNP) and the dependent (facial shape pre segmentation) variables using partial least squares regression (plsrregress from Matlab 2017b), and thus performed the canonical correlation analysis under a reduced model with residualized variables.

**Recruitment**

Two publicly available datasets, one from FaceBase and one from the ALSPAC project were used and did not involve any recruitment specific to this work. The other datasets used from PSU and IUPUI, and their characteristics, as mentioned above, were recruited over different studies and did not contain any specific selection or bias that might influence this work.

**Ethics oversight**

We have complied with all relevant ethical regulations for work with human participants and informed consent was obtained. Institutional review board (IRB) approval was obtained at each recruitment site and all participants gave their written informed consent prior to participation; for children, written consent was obtained from a parent or legal guardian. For the 3DFN sample, the following local ethics approvals were obtained: Pittsburgh, PA (PITT IRB PRO09060553 and R80405013); Seattle, WA (Seattle Children’s IRB 121107); Houston, TX (UT Health Committee for the Protection of Human Subjects HSC-DB-09-0508); and Iowa City, IA.
IA (University of Iowa Human Subjects Office IRB [200912764 and 200710721]. For the Penn State sample, the following local ethics approvals were obtained: Urbana-Champaign, IL (PSU IRB 13103); New York, NY (PSU IRB 45727); Cincinnati, OH (UC IRB 2015-3073); Twinsburg, OH (PSU IRB 2503); State College, PA (PSU IRB 44929 and 4320); Austin, TX (PSU IRB 44929); and San Antonio, TX (PSU IRB 1278). For the IUPUI sample, the following local ethics approvals were obtained: Indianapolis, IN and Twinsburg, OH (IUPUI IRB 1409306349). For the ALSPAC sample, approval was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committees. Consent for biological samples was collected in accordance with the Human Tissue Act (2004).

Note that full information on the approval of the study protocol must also be provided in the manuscript.