Integrative approaches generate insights into the architecture of non-syndromic cleft lip with or without cleft palate

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Summary

Non-syndromic cleft lip with or without cleft palate (nsCL/P) is a common congenital facial malformation with a multifactorial etiology. Genome-wide association studies (GWASs) have identified multiple genetic risk loci. However, functional interpretation of these loci is hampered by the underrepresentation in public resources of systematic functional maps representative of human embryonic facial development. To generate novel insights into the etiology of nsCL/P, we leveraged published GWAS data on nsCL/P as well as available chromatin modification and expression data on mid-facial development. Our analyses identified five novel risk loci, prioritized candidate target genes within associated regions, and highlighted distinct pathways. Furthermore, the results suggest the presence of distinct regulatory effects of nsCL/P risk variants throughout mid-facial development and shed light on its regulatory architecture. Our integrated data provide a platform to advance hypothesis-driven molecular investigations of nsCL/P and other human facial defects.

Introduction

Current research into the etiology of common disorders is focused on the identification of genetic susceptibility factors and the manner in which these risk variants interfere with biological function. Over the past decade, genome-wide association studies (GWASs) of common disorders have identified numerous risk loci. However, success in the translation of statistical associations from GWASs into functional mechanisms is only a very recent achievement. 1–6 A major driver of these advances has been the availability of large-scale genetic data and the systematic integration of genetic, transcriptional, epigenetic, and other -omics datasets from disease-relevant cell types and tissues. 7

Facial disorders rank among the most common birth defects worldwide and represent a substantial burden for affected individuals, their families, and healthcare systems. 6, 8 The most frequent facial disorder is non-syndromic cleft lip with or without cleft palate (nsCL/P). This condition has a global incidence of ~1 in 1,000 live births, 9 and is characterized by a multifactorial etiology that includes an overall genetic contribution of around 90%. 9–11 On an epidemiological level, nsCL/P is associated with an increased risk for adverse health outcomes. 12 However, this observation remains largely unexplained at both the clinical and molecular levels. To date, GWASs and other systematic approaches have identified at least 40 nsCL/P risk loci, 13–28 which explain up to 30% of the estimated heritability in European populations. 21 Despite these successes, functional dissection of the associated regions has been limited to only a few loci. 29–32 This is mainly attributable to the systematic underrepresentation of embryonic facial data in public resources such as ENCODE, 33 Roadmap Epigenome, 34 and GTEx. 35 To overcome this limitation, researchers have recently profiled multiple chromatin modifications in cell types and tissues of relevance to individual time points of mid-facial development, a process that is largely completed by week 10 of gestation (Figure 1A). These cell types and tissues include early human neural crest cells (hNCCs), 37 lineage-specific human cranial NCCs (cNCCs), 38 and embryonic mid-facial tissue samples encompassing the time period 4.5–10 weeks post-conception (craniofacial tissue [CT]; days 32–56 of gestation). 39 Previous studies have demonstrated a significant enrichment of nsCL/P-GWAS

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variants in active chromatin regions from both hNCCs and CT. To date, however, the fact that these datasets have been generated from differing sources has precluded the integrative analyses required for a comprehensive assessment of variant function at different time points of mid-facial development.

To generate novel insights into the etiology of nsCL/P, the present study leveraged both existing GWAS data on nsCL/P and epigenetic data on mid-facial development. The specific aims of the study were threefold (Figure S1). First, we generated one of the largest genome-wide genetic datasets for nsCL/P to date by combining three GWASs, which collectively encompassed European, Asian, and Latin American ethnicities. Using this resource, which we term MAiC (meta-analysis in clefting), we confirmed the vast majority of established risk regions and detected five novel loci (the strategy for identification of novel risk loci is described in the Supplemental Material and methods). To shed light on potential etiological overlaps between nsCL/P and other phenotypes, we then cross-referenced the lead variants at nsCL/P risk loci with GWAS data on >3,000 common traits and identified a set of loci with pleiotropic effects. Second, we compiled a comprehensive epigenetic map of mid-facial development through joint analyses of available data from hNCCs, cNCCs, and CT. This resource of chromatin segments

Figure 1. Human facial development and results of meta-analysis in clefting (MAiC)

(A) Schematic representation. The first phase of facial development (blue shading) is characterized by a substantial contribution of neural crest cells (NCCs): In early embryogenesis, NCCs arise in the ectoderm, undergo epithelial-to-mesenchymal transition, and begin to migrate from the dorsal neural tube. An NCC fraction (i.e., cranial NCCs) contribute to the pre-swellings of the face and populate the future frontonasal prominence as well as the first (purple) and second (green) pharyngeal arches. Subsequently, NCC-derived cells fuse to form those human facial structures that are finalized by the 10th week of embryogenesis.

(B) MAiC quantile-quantile plot. Observed statistical associations for non-syndromic cleft lip with/without cleft palate (nsCL/P) are plotted against the association statistics expected under the null hypothesis of no association. The contribution of different ethnicities in MAiC is shown using a pie chart.

(C) MAiC Manhattan plot. MAiC $-\log_{10}(p)$ association results are plotted along their chromosomal distribution. Blue and red lines indicate suggestive ($p < 10^{-5}$) and genome-wide ($p < 5 \times 10^{-8}$) significance, respectively. The lowest $p$ value was observed for rs55658222 ($p = 8.69 \times 10^{-63}$), located at 8q24.27 Novel risk loci are highlighted in green (lead variant plus variants in linkage disequilibrium [LD] $|r^2| \geq 0.6$). Gene names in subscript discriminate novel risk loci in situations where the respective chromosomal band is already listed among the 40 risk loci.
across mid-facial development serves as a platform for the interpretation of genetic findings for facial disorders and traits. Finally, we aimed to generate systematic insights into nsCL/P biology by combining MAiC and epigenetic data and then adding additional layers on gene expression in NCCs and global and local three-dimensional (3D) genomic interactions (i.e., topologically associated domains [TADs]), promoter-capture HiC [pCHi-C]). This approach revealed tissue- and time-point-specific regulatory effects at GWAS risk loci, prioritized candidate target genes, and highlighted distinct pathways. To our knowledge, the present report is the first to describe the systematic integration of large-scale summary statistics in nsCL/P and data on the cis-regulatory landscape across several stages of human mid-facial development.

Material and methods

GWAS meta-analysis MAiC

Cohort description

The meta-analysis included data from three previously published individual GWASs on nsCL/P (Bonn case-control GWAS cohort, GENEVA trio cohort, and POFC GWAS cohort). We included all nsCL/P summary statistics that were publicly accessible until June 2018. Data from the Bonn cohort were available in-house, while both the GENEVA (dbGaP: phs000094) and POFC (dbGaP: phs000774) datasets were downloaded from dbGaP upon approved data access, respectively. Previously conducted meta-analyses included combinations of two of these studies (Bonn and GENEVA GWAS cohort in Ludwig et al., 2012 [genotyped variants] and 2017 [imputed variants], GENEVA and POFC in Leslie et al.20). In the present study we combined the three GWAS cohorts to generate the largest nsCL/P meta-analysis to date. In accordance with previous studies, two meta-analyses were performed: (1) using all individuals with diverse population backgrounds (to increase statistical power by maximizing sample size; in the following termed as MAiC), and (2) using the European datasets only (MAiC_Euro) to reduce genetic heterogeneity based on population differences. Data quality control (QC) included the detection and removal of overlapping individuals, confirmation of ethnicity, and data re-analysis. We call this new dataset MAiC to provide a clear distinction from the previous individual studies and meta-analyses of sub-cohorts. Further details in cohort description and data QC can be found in the Supplemental information.

Statistical analyses

Statistical analyses were performed separately for case-control cohorts and case-parent trios, respectively. Imputed data were taken as provided by dbGaP (POFC) or generated as previously described (for Bonn and GENEVA), respectively, and best-guess genotypes were assigned based on a posteriori genotype probabilities of ≥0.6. In the case-control cohorts, GWAS was performed using logistic regression performed with SNPTEST and -method expected, by incorporating five (Bonn and GENEVA cohorts) and 18 (POFC cohort) dimensions of the multi-dimensional-scaling coordinates, respectively. For the case-parent trios, a transmission disequilibrium test (TDT) was performed on the best-guess genotypes. After data cleaning procedures (Supplemental information), we meta-analyzed the GWAS data of all four sub-cohorts (Bonn case-control, GENEVA case-parent trios, POFC case-control, and POFC case-parent trios) using METAL.

The final MAiC dataset (case-control plus case-parent trios) contained 6,825 individuals (including 3,946 affected; MAiC_Euro: 3,568 individuals including 1,517 affected; Table S1). The maximum genomic inflation factor was 1.051 (GENEVA) and 1.056 (POFC case-control) for MAiC and MAiC_Euro, respectively. All functional downstream analyses are based on MAiC because of largely increased statistical power. To estimate the single-nucleotide polymorphism (SNP)-based heritability (h²) for nsCL/P on the liability scale, we generated a European case-control-only dataset (Bonn, POFC, totaling 532 cases and 2,051 controls; Table S1) and performed linkage disequilibrium (LD) score regression as implemented in lddr.3 Sample and population prevalence were set to 0.21 and 0.001, respectively.

Gene-based and pathway analyses

Gene-based analyses in MAiC and MAiC_Euro were performed using MAGMA (v.1.06), implemented in FUMA. The input SNPs of MAiC were mapped to 17,911 protein-coding genes based to a distance of 0 kb upstream/downstream of the genes, resulting in a threshold of test-wide significance of p = 2.79 × 10⁻⁴⁶ (i.e., 0.05/17,911). To annotate known and novel nsCL/P risk loci in biological context, we investigated common expression patterns of the GWAS_TAD genes and their molecular functions (gene ontology [GO] terms, Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways) using FUMAs “GENE2FUNC” tool in (1) all GWAS_TAD genes, and (2) a subset of GWAS_TAD genes expressed in NCCs. This approach allows us to pinpoint risk loci or genes that are functionally involved in the same pathways or molecular processes and might be useful for gene prioritization.

Analysis of pleiotropic effects using the GWAS ATLAS

For each of the 45 lead SNPs in MAiC, association signals from large-scale genetic studies (including p value, effect size, and effect direction) were retrieved from the GWAS ATLAS. At time of analysis (November 2019), the database comprised 4,756 GWASs on 3,302 unique traits. Notably, the unique traits are split into 28 domains, of which we combined two (environment, activities) into one domain to reduce redundancy. All significant SNP-trait associations at p < 0.05 were considered, and this number was corrected for the number of GWASs and loci in the analysis.

Epigenetic datasets for mid-facial development

Identification of datasets relevant to mid-facial development

Human cell-type- and developmental-stage-specific data for mid-facial development are underrepresented (or not represented at all) in large consortia data such as ENCODE.35 However, available data in the Gene Expression Omnibus (GEO) covered mid-facial development from (1) early stages (hNCCs, accessed through GEO: GSE28874), (2) differentiated human cNCCs (accessed through GEO: GSE70751), and (3) embryonic craniofacial human tissue of different Carnegie stages (CS) (accessed through GEO: GSE97752). In each of these datasets, analyses of chromatin modifications were performed using chromatin immunoprecipitation followed by sequencing (chromatin immunoprecipitation sequencing [ChIP-seq]) or are available as imputed datasets. Detailed information including antibodies used in these studies is shown in Table S3 and in the Supplemental information. For hNCCs and cNCCs, ChIP-seq had been performed for chromatin modifications H3K27ac, H3K4me1, H3K4me3, and H3K27me3. In CT, for samples of CS13–CS17, ChIP-seq was performed for H3K27ac, H3K4me1, H3K4me3, H3K27me3, and H3K36me3 (Table S4), and data for H3K9me3 were imputed. For CS20 and...
10 wpc, H3K27ac3 ChIP-seq data were experimentally derived; all other marks were imputed (Table S3).

**Data processing**

For hNCCs and cNCCs, raw data were available in fastq format. A description of data QC is given in Rada-Iglesias et al. and Prescott et al., respectively. ChIP-seq data from craniofacial data in Wilderman et al. comprise processed formats, including imputed signals, peaks, and segmentation data. In order to ensure comparability among the three data sources, computational processing of ChIP-seq data as published in Wilderman et al. (QC, alignment, peak calling, epigenetic imputation, chromatin segmentation) was adopted in the hNCC/cNCC bioinformatics pipeline, as described in the Supplemental information and Table S3.

**Chromatin imputation and segmentation**

To obtain uniform datasets, chromatin imputation followed by chromatin state segmentation was performed. First, H3K9me3 and H3K36me3 marks in hNCCs/cNCCs were imputed using ChromImpute (v.1.0.1.46 based on 127 cell types from the Roadmap Epigenome Project. Imputed hNCC/cNCC signal files for each individual chromosome and each chromatin mark were binarized, and segmentation was performed using the core+K27ac 18-state chromatin model provided by Roadmap with ChromHMM to predict 18 chromatin states. Because of the low number of chromatin marks measured in the NCC samples, epigenetic imputation issues, and the higher risk of batch effect between hNCCs, cNCCs, and CT, we adopted a robust strategy and condensed the 18 generated states into eight states, based on Roadmap definition: three active states (transcription starting sites [TSS], transcribed sites, and enhancers [Enh]), one bivalent state (Poised Enh/bivalent TSS), three repressed states (Heterochromatin, Repressed PolyComb sites, Zinc finger genes/Repeats), and one quiescent state (Quies). Potential batch effects were analyzed using principal-component analysis (PCA) and hierarchical clustering of Pearson correlation coefficients.

**Other datasets**

To identify genome-wide regulatory genomic units, we used TADs from human embryonic stem cells (hESCs) (H1 cell line) as provided by the Ren Lab. Protein-coding genes were extracted from UCSC genome browser (hg19) and were mapped to TADs using positional information. TADs containing an nsCL/P risk locus were defined as GWASTAD region. Based on previous evidence for complex regulatory interactions within one TAD, we considered all genes from the GWASTAD region as potential candidate genes for downstream effects of the associated variants in the r^2 ≥ 0.6 region. Expression data from NCCs (two replicates of day11hNCC [GEO: GSE121428] and three replicates of passage2hNCC [GEO: GSE108521]) were retrieved from Laugsch et al. (GEO: GSE108522). For the comparison of genes in TADs of nsCL/P risk loci and genes expressed in NCCs, we used the average RNA-seq Fragments Per Kilobase Million (FPKM) across five samples. To identify functional links between different regulatory features (e.g., DNA-DNA interactions of enhancers and TSS) at specific risk loci, we accessed pCHi-C cis-interaction data collected in hESCs (GEO: GSE86821). 51

**Translation of genetic associations into tissue- and time-point-specific regulatory effects at a systematic level**

**Enrichment analyses using GREGOR**

Based on chromatin segments obtained from hNCCs, cNCCs, and CT, we used GREGOR (Genomic Regulatory Elements and GWAS Overlap Algorithm) 51 to evaluate the enrichment of significant SNPs from the MAiC data in the available regulatory features (i.e., eight predicted chromatin states). As described in the Supplemental information, a set of samples from the Roadmap Epigenomics project (comprising both fetal and adult tissue samples) was selected as an independent dataset for comparison. As input, we used MAiC nsCL/P variants with p ≤ 0.001 without additional variants in LD (n = 22,999); this threshold was selected to balance between adequate statistical power and true-positive association signals.

**CT- and NCC-specific active chromatin sites**

To examine specific effects in either NCCs or CT, we filtered in the chromatin segmentation datasets for active chromatin sites (TSS, Enhancer or transcribed sites) in NCCs that are repressed/quiet (Quiescent, Biv_TSS_pois_enh, RepPrFC, Heterochromatin) in CT and vice versa. For robust observations, we only trust in a chromatin state if it is present in both NCC samples (hNCCs, cNCCs) or in five of the six CT (CS13, CS14, CS15, CS17, CS20, 10wpc) samples. To account for biases in length associated with batch effects, active sites were only retained if they had a distance of ≥500 bp to any chromatin segment of opposite active status in the other cell system/tissue. In the following, we combined the specific active chromatin sites with MAiC associations and TAD data to filter for TADs with high density of strong associated genetic variants (pMAiC ≤ 5 × 10^-5) in specific active chromatin sites at new and known nsCL/P GWAS risk loci.

**Characterization of nsCL/P risk variants and candidate gene prioritization in context of epigenetic mid-facial timeline**

For comprehensive insights in regulatory mechanisms at nsCL/P risk loci, we finally integrated all available genetic and functional data (MAiC associations, GWASTAD- and r^2 ≥ 0.6-region boundaries, NCC- and CT-specific active chromatin sites, chromatin segmentation tracks, and pCHi-C cis interactions). Based on this approach, we attempt to prioritize genetic variants with regulatory effect and potential downstream target genes and to detect relevant regulatory elements specific for the early (hNCC/cNCC) or later mid-facial development (CT).

**Results**

**MAiC identifies five novel risk loci**

The MAiC dataset was generated by combining GWAS data from three previous studies (Bonn, GENEVA, POFC), following the exclusion of overlapping individuals and extensive QC. The final dataset comprised 1,247 nsCL/P cases, 2,879 controls, and 2,699 case-parent trios of multiple ethnicities, and ~7.74 million SNPs. The p value distribution was consistent with a multifactorial inheritance (Figure 1B; lambda = 1.07). A set of 1,375 SNPs achieved genome-wide significance (p < 5 × 10^-8; Figure 1C). Analysis of established nsCL/P risk loci in MAiC revealed genome-wide significant SNPs at 25 of the 40 regions. These 25 regions comprised 22/26 loci that were previously identified in GWASs based on largely European samples and 3/14 loci reported in individuals from the Chinese population. At all other nsCL/P risk loci (n = 15), nominal significance (p < 0.05) was observed for individual
variants that were in strong LD (D’ > 0.8) with the respective lead SNP (Table S2).

Importantly, the MAiC analyses also identified five novel risk loci (p < 5 x 10⁻⁵), thus increasing the number of identified nsCL/P GWAS risk loci to 45. These novel loci were located at chromosomes 1p36.13 (sentinel variant rs34746930), 5p12 (rs60107710), 5q13.1 (rs6449957), 7p21.1 (rs62453366), and 20q13.12 (rs3091552; Table 1). Consistent with previous findings on risk variants for nsCL/P and other complex traits, these lead variants map to non-coding regions that are adjacent to candidate genes with functions during facial development, such as CAPZB52 and NBL153 (both at 1p36) and EYA254 (at 20q13; Supplemental text; Figures S2–S6). To identify population-specific effects, a sub-analysis was performed in individuals from Central Europe (MAiC euro; n = 562 cases, 2,051 controls, and 955 case-parent trios). No additional risk loci were identified at the level of genome-wide significance (Figure S7; Table S2). Using this European case-control cohort and LD score regression, SNP-based heritability was estimated as h² = 28% ± 0.1%. This confirmed previous heritability estimates obtained using the Bonn cohort only.21

Gene-based analyses suggest nsCL/P candidate genes outside of GWAS risk loci
Using MAiC summary statistics and MAGMA,46 gene-based analyses yielded 1,357 genes with nominal significance (p < 0.05; Figure S8A). A total of 25 genes reached test-wide significance (p < 2.79 x 10⁻⁶; Table S6). Of these, 23 map to known GWAS risk loci. For some of these 23 genes, functional evidence strongly supports their involvement in nsCL/P (e.g., IRF6, TP6356). This analysis also suggested novel candidate genes at GWAS risk loci, such as ARID3B. In mice, the gene Arid3b is expressed in cranial mesenchyme structures and has been shown to interact with Mycn, which is encoded by a strong candidate gene with classical inheritance pattern in multigenerational families (four genes), or (3) an enrichment of rare coding variants (five genes).50 Of these, 12 genes were present in the analysis set. Two of these 12 genes approached test-wide significance: PRTG (p = 8.44 x 10⁻⁵) and CTNND1 (p = 2.17 x 10⁻⁵; Table S8). These observations indicate that in at least a subset of genes, both common and rare variations, contribute to nsCL/P.

Genes located in TAD regions of nsCL/P GWAS loci are enriched in developmental pathways
Accumulating evidence suggests that most regulatory interactions occur within TAD modules.63,64 Therefore, genes located within TADs represent candidates for the downstream effects of the associated common risk variants. To identify molecular processes of relevance to nsCL/P, for each of the 45 risk loci, GWAS-TAD regions were defined, based on the extent of the respective TAD in hESC data.65 In total, 407 genes were identified within the respective TADs (GWAS-TAD genes, range 1 to 29 genes per locus; Table S9). Enrichment analysis using MAGMA yielded test-wide significant (p_adj ≤ 0.05) results for 287 GO terms (Table S10). The most significant enrichments were observed for “tissue development” (p_adj = 8.34 x 10⁻⁶), “epithelium development” (p_adj = 8.82 x 10⁻⁶); and “appendage development” (p_adj = 7.92 x 10⁻⁶; Figure S10). Together with additional significant terms, such as “embryo development,” “tube development,” and “ear development,” these observations suggest the existence of common pathways for nsCL/P and other processes of organogenesis during embryonic development.

We then prioritized genes expressed in NCCs by adding available RNA sequencing (RNA-seq) data from hNCCs.66 In total, 240 of the 407 GWAS-TAD genes were expressed in NCCs, with strong expression being observed for a subset of 12 genes (≥ 200 fragments per kilobase mapped; Table S9). Of these, at least two have been previously implicated in NCC migration processes (CAPZB, TP63). These 240 NCC-expressed genes showed a substantial overlap in significant GO terms compared with the analysis of all 407 GWAS-TAD genes (233 out of 287 pathways; Figure 2A; Table S11). Of those 233 pathways, 157 pathways showed stronger enrichment in the subset of NCC-expressed GWAS-TAD genes, the strongest of which represent cellular processes (Figure S10; Table S12). Among pathways that were exclusive to GWAS-TAD genes expressed in NCCs (n = 106), both regulatory processes and metabolic pathways were enriched. In contrast, pathways specific to GWAS-TAD genes that were not expressed in NCCs (n = 54) included “keratinocyte proliferation” and “epidermis development,” a finding that is consistent with the substantial contribution of the epithelial lineage to nsCL/P.56

We next addressed the potential etiological overlap between nsCL/P and other common phenotypes that might contribute to the adverse health outcomes observed in nsCL/P. We retrieved association signals for each of the 45 lead SNPs in MAiC from large-scale genetic studies, using the GWAS ATLAS.47 At the time of analysis (November 22, 2019), this resource comprised 4,756 GWASs on 3,302 unique traits. While all of the 45 variants were available in the atlas, only 19 showed at least one significant SNP-trait association when corrected for the number of GWASs and...
loci (p < 2.33 × 10^-7; overall number: n = 219; Table S13). These associations reflect 35 collapsed traits across 12 domains, including height, bone mineral density, hair color, and body mass index (Table S14). Eighteen traits showed associations with at least two distinct nsCL/P risk loci. Interestingly, for some traits, the direction of effect differed between individual loci (e.g., height and bone mineral density), while for other traits, the direction of effect was consistent (e.g., hypothyroidism, glomerular filtration rate, and hair color; Figure 2B).

**NsCL/P-associated variants are enriched in multiple chromatin states of mid-facial development**

Recent analyses in human embryonic CT demonstrated both a significant enrichment of lead SNPs from earlier nsCL/P GWASs in active enhancers and the presence of mid-facial specific regulatory elements. To extend this work, we incorporated data from two NCC states in order to generate a unified mid-facial development resource of chromatin modifications (Figure S1). We retrieved data on ChIP-seq from hNCCs and cNCCs and applied the data analysis pipeline used by previous authors for computational analyses of ChIP-seq data from CT. We observed strong inter-sample correlations between chromatin mark and developmental stage (Figures S11 and S12). The integration of 127 non-facial samples from Roadmap revealed local clustering of NCCs and CT along a hierarchical axis comprising hESCs, induced pluripotent stem cells (iPSCs), and iPSC-derived cells (Figure S13). Here, the most tissue-specific pattern was observed for H3K27ac (Figure S14). Similar to a previous finding for CT, non-facial fetal tissue samples (such as brain, kidney, and lung) clustered distinctly from NCCs (Figure S14), thus emphasizing the limited utility of many public resources for the interpretation of genetic findings in facial disorders.

Next, we generated robust chromatin segments in NCCs using ChromHMM. Together with segmentation data from CT and Roadmap, chromatin segments were condensed to eight categories in order to increase the robustness of the subsequent analyses (Figure S15; Table S15). We then analyzed the positional overlap of all variants with pMAIC < 0.001 in the eight chromatin states across NCCs and CT (SNP0.001_nsCL/P n = 22,999), and compared this to a matched set of non-associated SNPs (SNPcontrol_nsCL/P p > 0.1). The results showed that 23% of the nsCL/P variants (SNP0.001_nsCL/P) mapped to active chromatin states, while 14% mapped to either bivalent or repressed chromatin states (Figure 3A). This enrichment was significantly higher compared to the control SNPs, where 16% and 11% of variants mapped to active, or to bivalent/repressed, chromatin states, respectively (p < 10^-16; Fisher’s exact test).

To delineate associations of specific chromatin states along the time series, enrichment was tested using GREGOR. For each of the two SNP sets, every hNCC/cNCC/CT sample was tested, together with 11 randomly selected Roadmap samples (both fetal and adult). A significant enrichment for SNP0.001_nsCL/P was observed in most of the samples/chromatin states (Figure 3B; Table S16), as compared to SNPcontrol_nsCL/P (Figure S16; Table S17). While the fold enrichment (FE) was similar for NCCs and CT in six of the eight chromatin states (such as those related to active transcription; Figures 4A–4D; Figure S17), considerable differences in enrichment between NCC and CT samples were observed in chromatin states “active enhancers” and “poised enhancers/bivalent TSS.” In both states, NCCs displayed a stronger enrichment than CT samples. For enhancers, the mean FE (FEMean) in NCCs was 1.64 (pMean = 4.36 × 10^-86), average of pGREGOR), compared with FEMean = 1.43 in CT (pMean = 8.09 × 10^-22). For “poised enhancers/bivalent TSS,” the corresponding values were FEMean = 1.65, pMean = 3.39 × 10^-20 in NCCs, compared with FEMean = 1.39, pMean = 4.74 × 10^-4 in CT. These results may have been driven in part by the heterogeneous composition of the CT samples. However, the specific enrichment pattern observed in two out of eight chromatin states suggests a distinct biological underpinning. Overall, these data confirmed previous findings of an over-representation of nsCL/P lead variants in enhancer marks and extended this enrichment toward additional common variants and annotations.

**A subset of nsCL/P-associated SNPs show distinct regulatory effects**

To extend the investigation of the contribution of regions with differing regulatory profiles in NCCs and CT, we
created genome-wide maps of active chromatin sites for both NCCs and CT. A total of 9,897 regions (encompassing 26.67 Mb) with active chromatin states in NCCs (TSS, enhancer or transcribed sites) were inactive in CT (quiescent, repressed, or bivalent; termed NCC-specific active sites). Similarly, 6,189 regions (29.37 Mb) were active in CT but inactive in NCCs (CT-specific active sites). The integration of MAiC association data revealed 62,084 genetic variants that map in NCC-specific active sites. Of these, 4,022 had $p_{\text{MAiC}} < 0.05$. Similarly, 72,556 variants (4,834 of which had $p_{\text{MAiC}} < 0.05$) mapped to CT-specific active sites. In each of the groups of NCC-specific and CT-specific active sites, the $p$ value distribution differed significantly from that expected, with a significant enrichment of association signals being observed at the lower tail of the distribution (Figure 5A).

Filtering for the subset of SNPs with $p_{\text{MAiC}} < 5 \times 10^{-5}$ identified 112 SNPs that mapped to either NCC-specific (51 variants), or CT-specific active regions (61 variants; Table S18). These were distributed over 39 TADs, which encompassed both known nsCL/P risk loci (n = 19; e.g., chromosomes 1p22 [Figure S18] and 2p24.2 [Figure S19]).
regions with suggestive evidence for association \((n = 20); \text{ e.g., chromosome 4p13 [Figure S20]). Interestingly, at six loci (e.g., chromosomes 1q32.1 [Figure S21] and 15q24.1 [Figure S22]), at least two associated variants in LD were located in different specific elements (Table S19). This represents a significantly higher enrichment than expected and suggests that individual variants of risk haplotypes might affect the regulatory architecture at different stages of craniofacial development (Figure 5B).}

Finally, we assessed how novel hypotheses on nsCL/P pathogenesis can be generated from the systematic integration of data concerning: (1) statistical associations (MAiC), (2) chromatin modifications over time (mid-facial time-series), and (3) pCHi-C cis-interactions.\(^{31}\) Examples from two loci are described here. First, at 5q13 PIK3R1, the lead variant \(\text{rs6449957} \quad \text{pMAiC} = 6.59 \times 10^{-10}\) is located within an active region upstream of \(\text{PIK3R1}\). This region shows evidence of being transcribed but lacks any RefSeq annotation, which might point toward a transcribed enhancer or an as-yet-undetected transcript. PCHi-C data indicate \(\text{cis}\) interactions with \(\text{PIK3R1}\) and \(\text{MAST4}\), both of which are expressed in hNCCs. In addition, another variant in strong LD \(\text{rs921792} \quad \text{pMAiC} = 1.17 \times 10^{-5}\) maps to a putative enhancer that is detected in both NCCs and CT (Figure 5C). As a second example, at 13q32.2 (lead variant \(\text{rs2763990} \quad \text{pMAiC} = 3.03 \times 10^{-6}\), intronic in \(\text{CLYBL}\) ), interactions were observed between the region around the lead variant and the genes \(\text{ZIC2, ZIC5, and GGACT}\). While some variants (including \(\text{rs2763934}\) with \(\text{pMAiC} = 6.53 \times 10^{-7}\)) map to a craniofacial active element near the \(\text{CLYBL}\) gene promoter, additional variants (including \(\text{rs4525350}\) with \(\text{pMAiC} = 6.39 \times 10^{-6}\)) map to several more distantly located NCC-specific enhancers. Based on pCHi-C data, our data indicate that in NCCs, risk variants might affect \(\text{ZIC2}\) and \(\text{ZIC5}\) expression. This hypothesis is further supported by the finding of active transcription sites in NCCs and a bivalent state in embryonic and adult tissues. A plausible hypothesis is that, at later time points of development, additional variants mapping to other enhancer elements act on \(\text{GGACT}\), as suggested by the presence of transcribed sites in CT. Notably, the transcript region of \(\text{CLYBL}\) itself has limited evidence for active transcription across all analyzed stages of mid-facial development, despite the presence of some active marks in the promoter region (Figure 5D).}

At other loci, our data provide evidence for the presence of tissue-specific gene isoforms (e.g., 4p13-locus; Figure S20), or a second, novel candidate gene at previously reported loci. For example, at chromosome 1p22, our data suggest that the previously identified gene \(\text{ARHGAP29}\)\(^{16}\) is a target gene with CT-specific expression and highlight \(\text{ABCD3}\) as novel candidate gene (Figure S18). The data also suggest complex promoter-promoter interactions involving all genes at this locus (\(\text{ARHGAP29, ABCD3, and ABCA4}\)). Interestingly, the MAiC top-associated variant at 1p22 \(\text{rs352988667} \quad \text{pMAiC} = 6.86 \times 10^{-16}\) has putative enhancer function and maps to the “E2” element, whose functional role in nsCL/P was confirmed in previous research.\(^{32}\) At another locus (1q32.1), we found that \(\text{SERTAD4}\) is a CT-specific target gene, while the established causal gene \(\text{IRF6}\) was marked as bivalent, which...
is consistent with its established activity in epithelial tissue \textsuperscript{55,68} (Figure S21). Taken together, these results will inform future functional studies into nsCL/P and underscore the importance of the thorough genomic annotation of relevant cells and tissues.

Discussion

Here, we report on a data-driven approach that generated novel insights into the etiology of nsCL/P. At the genetic level, we identified five novel risk loci via the large-scale meta-analysis of common genetic variation. This large genome-wide resource empowered systematic analyses at the gene and pathway levels and implicated novel molecular players in nsCL/P. Our analysis of pleiotropic effects on other common traits revealed a substantial positional overlap with traits such as height and bone mineral density. At some loci, associated variants showed opposite directions of effect, which indicates their contribution to distinct pathways. We have provided examples of how this resource is useful in terms of translating statistical associations into biological insights and illustrated its potential for further analyses of facial disorders and traits.

While our results are based on a multiethnic cohort, this still comprises a substantial contribution from the European population. Still, we captured associations at all loci that had been previously reported in distant ethnicities, such as the Chinese population. Although these observations suggest that nsCL/P might show less locus heterogeneity than is the case for other common diseases, allelic heterogeneity is likely to contribute in part to the lack of replication observed at some loci in previous studies. Also, the integration of genetic and chromatin segmentation data might have been biased by the European background of both the genetic and epigenetic maps. Despite some initial evidence that methylation patterns show population-specific components \textsuperscript{69,70}, few studies to date have performed systematic analyses of how maps of chromatin accessibility (in particular in mid-facial development) vary across populations. Future studies are required to determine whether population-specific risk variants from non-European populations show differing enrichment patterns from those observed in the present study and to identify additional pleiotropic effects that are present at other risk haplotypes in other populations. Importantly, to address these issues, future meta-analysis should also include recent GWAS data (e.g., from Sub-Saharan Africans \textsuperscript{71} and Colombians \textsuperscript{72}). In addition, our analyses were performed for nsCL/P as the central trait. Previous studies have generated evidence of an (albeit incomplete) etiological overlap between the various nsCL/P subtypes (e.g., cleft lip, and cleft lip with cleft palate) and the genetic heterogeneity of other types of orofacial clefting (e.g., cleft palate only). Application of our integrative approach to the investigation of cleft subtypes will facilitate understanding of their individual etiologies, an issue that was beyond the scope of the present study.

One major feature of our approach was that it combined previous individual data into one joint map of epigenetic chromatin segments of NCCs and CT. This will be highly useful in terms of the future interpretation of associations in facial disorders/traits. However, due to limited availability of datasets from other cell types, such as human embryonic epithelium, this map does not comprehensively capture all biological contributors to human craniofacial development. Furthermore, our joint analysis of the different CT stages may have overlooked some effects within single stages of CT. Nonetheless, the data obtained at individual loci add to increasing evidence that for nsCL/P development, risk loci have a complex regulatory...
architecture, and several genes at single loci might be of relevance across the different time points of craniofacial development. Notably, several of the genes prioritized by our systematic approach have obtained independent support by other studies, for instance clefting syndromes (e.g., TP63, EEC syndrome\textsuperscript{75}), resequencing studies (e.g., ARHGAP29\textsuperscript{76,77} and IRF6\textsuperscript{25}), or experimental evidence (e.g., PAX7\textsuperscript{78}). While we here focused on an \textit{in silico} approach, we hope that the results will empower further experimental investigations of specific risk variants that were highlighted among the set of associated variants. Using the joint pipeline, we will continue to update our resource as chromatin marks become available from additional human tissues and/or cell systems of relevance to mid-facial development. In addition, the map will be refined through the use of single-cell technologies in order to resolve the issue of tissue heterogeneity encountered in the present study. Finally, the integration of other layers of genetic information, such as rare variants identified by whole-exome or -genome sequencing in cleft cohorts, will further increase our understanding of the etiology of craniofacial development and disease.\textsuperscript{61,79}
Data and code availability

Original data for genetic and functional analyses in the paper is available as follows: dbGaP (dbGaP: phs000094 and phs000774), GEO (GEO: GSE28874, GSE70751, and GSE97752), and Zenodo (DOI 10.5281/zenodo.3724148). The NCC- and CT-specific active sites generated during this study are available at Zenodo (DOI 10.5281/zenodo.3911187).

Supplemental information

Supplemental information can be found online at https://doi.org/10.1016/j.xhgg.2021.100038.

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Declaration of interests

The authors declare no competing interests.

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Web resources

| ANNOVAR, https://wgdag.org/software/9-annovar |
| Bowtie2, http://bowtie-bio.sourceforge.net/bowtie2/index.shtml |
| ChromHMM, http://compbio.mit.edu/ChromHMM/ |
| ChromImpute, http://www.biochem.ucl.ac.uk/labs/ernst/ChromImpute/ |
| core 15-state chromatin model, https://egg2.wustl.edu/roadmap/data/byFileType/chromatinSegmentations/ChmmModels/ |
| FastQC, https://www.bioinformatics.babraham.ac.uk/projects/fastqc |
| FUMA, https://fuma.ctglab.nl/ |
| GREGOR, http://csg.sph.umich.edu/GREGOR/ |
| GTEx, https://www.gtexportal.org/home/ |
| GWAS Atlas, https://atlas.ctglab.nl/ |
| Hi-C data from Bing Ren Lab, http://chromosome.sdsuc.edu/mouse/hi-c/download.html |
| IMPUTE2, http://mathgen.stats.ox.ac.uk/impute/impute_v2.html |
| KING: Kinship-based Inference for GWAS, https://www.kingrelatedness.com/ |
| LDLink, https://ldlink.nci.nih.gov/?tab=home |
| LDSTR, http://ldstr.broadinstitute.org/ldhub/ |
| MACS2, https://github.com/macs3-project/MACS |
| MAGMA, https://ctg.cncr.nl/software/magmA |
| METAL, http://csg.sph.umich.edu/abecasis/metal/ |
| MSigDB, https://www.gsea-msigdb.org/gsea/msigdb/index.jsp |
| PhantomPeakQualTools, https://github.com/kundajelab/phantompeakqualtools |
| Roadmap, https://egg2.wustl.edu/roadmap/data/byFileType/signal/consolidated/macs2signal/pval/ |
| UCSC, http://genome.ucsc.edu/ |

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