Comprehensive identification and analysis of human accelerated regulatory DNA

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It has long been hypothesized that changes in gene regulation have played an important role in human evolution, but regulatory DNA has been much more difficult to study compared with protein-coding regions. Recent large-scale studies have created genome-scale catalogs of DNase I hypersensitive sites (DHSs), which demark potentially functional regulatory DNA. To better define regulatory DNA that has been subject to human-specific adaptive evolution, we performed comprehensive evolutionary and population genetics analyses on over 18 million DHSs discovered in 130 cell types. We identified 524 DHSs that are conserved in nonhuman primates but accelerated in the human lineage (haDHSs), and estimate that 70% of substitutions in haDHSs are attributable to positive selection. Through extensive computational and experimental analyses, we demonstrate that haDHSs are often active in brain or neuronal cell types; play an important role in regulating the expression of developmentally important genes, including many transcription factors such as SOX6, POU5F2, and HOX genes; and identify striking examples of adaptive regulatory evolution that may have contributed to human-specific phenotypes. More generally, our results reveal new insights into conserved and adaptive regulatory DNA in humans and refine the set of genomic substrates that distinguish humans from their closest living primate relatives.

[Supplemental material is available for this article.]

A number of traits distinguish humans from our closest primate relatives, including bipedalism, increased cognition, and complex language and social systems (for review, see O’Bleness et al. 2012). To date, the genetic basis of human-specific phenotypes remains largely unknown, complicated by the difficulties in distinguishing between phenotypically significant and benign variation. Thus, evolutionary changes in protein-coding sequences have received considerable attention, as the phenotypic consequences of these mutations have historically been easier to interpret (Clark et al. 2003; Stedman et al. 2004; Chimpanzee Sequencing and Analysis Consortium 2005; Nielsen et al. 2005; Arbiza et al. 2006; Dennis et al. 2012; Sudmant et al. 2013). Although protein-coding evolution has clearly played a role in human evolution, proteins account for only ~1.5% of the human genome, most of which exhibit high sequence similarity between humans and chimpanzees (Chimpanzee Sequencing and Analysis Consortium 2005). However, between ~2.5% and 15% of the human genome is estimated to be functionally constrained (Chinwalla et al. 2002; Lunter et al. 2006; Asthana et al. 2007; Meader et al. 2010; Ponting and Hardison 2011). Thus, the mutational target size of noncoding DNA is considerably larger than protein-coding sequences, suggesting that regulatory DNA is also an important substrate of evolutionary change, as originally proposed four decades ago (Britten and Davidson 1969; King and Wilson 1975). In some cases, detailed studies of individual genes have revealed human-specific regulatory evolution, such as in FOXP2, which is thought to have influenced traits related to speech and language in humans (Enard et al. 2002).

Nonetheless, interpreting patterns of interspecific divergence and intraspecific polymorphism in noncoding DNA has been considerably more challenging compared with those of protein-coding sequences. An elegant and powerful way to identify evolutionary changes in noncoding DNA of potential significance, originally described by Pollard et al. (2006b) and extensively used thereafter (Pollard et al. 2006a,b; Prabhakar et al. 2006; Kim and Pritchard 2007; Bush and Lahn 2008; McLean et al. 2010; Lindblad-Toh et al. 2011; Pertsemlidis et al. 2011), focuses on the discovery of sequences that are rapidly evolving or lost on the human lineage but that are otherwise phylogenetically conserved and thus likely functional. This approach has led to the discovery of several regions with species-specific enhancer activity (Prabhakar et al. 2008; Capra et al. 2013; Kamm et al. 2013), as well as human-specific deletion of regulatory DNA (McLean et al. 2011).

However, phylogenetic conservation is an imperfect proxy for function, particularly for noncoding regulatory sequences that can exhibit significantly high rates of turnover (Dermitzakis and Clark 2002; Wray et al. 2003; Villar et al. 2014). To more directly identify regulatory DNA, recent studies such as the ENCODE (The ENCODE Project Consortium 2012) and Roadmap Epigenomics Projects (Bernstein et al. 2010) have created genome-scale maps of DNase I hypersensitive sites (DHSs) in a large number of cell types. DNase I preferentially cleaves regions of open and active DNA, making it a powerful assay to identify regulatory elements, regardless of their specific function (Galas and Schmitz 1978; Dorschner et al. 2004). Although high-resolution maps of DHSs now exist, not

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all experimentally defined regulatory elements are expected to be functionally or phenotypically significant (Eddy 2012; Doolittle 2013; Graur et al. 2013; Niu and Jiang 2013).

Thus, we hypothesized that the synergistic combination of comparative and functional genomics would facilitate the high-resolution identification of conserved and human accelerated regulatory sequences. Here we describe the genome-wide architecture and characteristics of 113,577 DHSs that are conserved in primates and 524 DHSs that exhibit significantly accelerated rates of evolution in the human lineage (haDHSs). We estimate that ~70% of substitutions within haDHSs are attributable to positive selection; we experimentally validated a large number of elements; and we perform extensive bioinformatics analyses that integrate information across multiple functional genomics data sets to better understand the functional and biological characteristics of haDHSs.

**Results**

**Framework for identifying conserved and human accelerated regulatory DNA**

To identify human accelerated regulatory DNA, we leveraged experimentally defined maps of DHSs from 130 cell types identified in the ENCODE and Roadmap Epigenomics Projects (Supplemental Table 1). After merging DHSs across cell types into 2,093,197 distinct loci (median size = 290 bp, SD = 159 bp), we used a whole-genome alignment of six primates from the EPO pipeline (Paten et al. 2008) to obtain separate alignments for each DHS, using strict filtering criteria for alignment quality. We performed two likelihood ratio tests to distinguish between DHSs that are evolving neutrally, are conserved among primates, or are conserved among primates but accelerated in the human lineage (haDHSs). Specifically, we used a maximum likelihood test (Pollard et al. 2010) to first identify 113,577 DHSs that exhibit significant evolutionary constraint across primates, which manifest as regions of low sequence divergence compared with carefully defined putatively neutral flanking sequence (FDR = 0.01) (Fig. 1). Next, for DHSs that are conserved in primates, we performed a second likelihood ratio test (Pollard et al. 2010) and identified 524 regulatory sequences that have experienced a significant acceleration of evolution in the human lineage and therefore exhibit an excess of human-specific substitutions (FDR = 0.05) (Fig. 1; Supplemental Table 2). Importantly, to avoid biasing ourselves against identifying human acceleration, we excluded the human sequence in the first test for conservation.

**Characteristics of primate conserved regulatory DNA**

We first characterized the set of DHSs conserved across primates. Approximately 93% of conserved DHSs overlap a phastCons conserved element, but many also contain short segments of less conserved sequence, making them overall less conserved than those identified by phastCons (Fig. 2A). We hypothesize that these less conserved sequences interspersed within DHSs may facilitate the rapid acquisition of novel transcription factor binding sites, as these regions are already actionable (i.e., accessible to proteins) and poised to evolve new functions compared with nonconserved sequences outside of DHSs.

Patterns of conservation varied significantly across cell type category (Kruskal–Wallis test; $P = 5.08 \times 10^{-8}$; Methods) (Fig. 2B), ranging from 5.0% of DHSs in chronic lymphocyte leukemia cells to 20.4% in fetal brain cells. DHSs active in fetal cell types showed the highest levels of conservation, consistent with the observation that gene regulation in developmental pathways is highly conserved (Lowe et al. 2011). Conversely, DHSs in malignant cell types exhibited the fewest conserved DHSs, which may reflect ectopic...
activation of chromatin (Vernot et al. 2012). These patterns are also observed in cell-type-specific DHSs (Supplemental Fig. 1a).

Genomic landscape of human accelerated regulatory DNA

We next investigated the set of haDHSs. Overall, these elements have evolved at approximately four times the neutral rate in the human lineage, while other primate branches have evolved at less than half of the neutral rate (Fig. 3A). In total, 70 haDHSs overlap previously identified human accelerated elements (HAEs) (Pollard et al. 2006b; Prabhakar et al. 2006; Bush and Lahn 2008; Lindblad-Toh et al. 2011), which is highly significant (permutation $P < 1 \times 10^{-5}$) (Fig. 3B). Thus, by focusing on experimentally defined regulatory DNA, we identify 454 novel loci that show accelerated rates of evolution in the human lineage, increasing the set of 1621 merged HAEs by 28%. The number of cell types each haDHS was active in varied substantially (Supplemental Fig. 2). Notably, 64% (337) of haDHSs were identified in at least one brain or neural cell type, and 88.5% (464) were active at least one developing fetal tissue.

In comparison to conserved nonaccelerated DHSs, haDHSs are significantly enriched in noncoding regions ($P = 1.16 \times 10^{-7}$, hypergeometric test) (Fig. 3C). These data are consistent with the hypothesis that noncoding regions are more free to evolve and acquire new functions. Furthermore, we observed eight regions where four or more haDHSs were clustered within a 1-Mb window, suggesting coordinated changes in multiple regulatory elements (Fig. 3D). For instance, TENM3, which is required for establishing neuronal connections in vertebrate retinal ganglion cells (Antinucci et al. 2013; Merlin et al. 2013), is the nearest gene to five haDHSs, four of which are active in retinal pigment epithelial cells (Fig. 3D, inset).

Adaptive evolution is the primary determinant of rate acceleration in haDHSs

Human acceleration can result from both adaptive and nonadaptive forces (Haygood et al. 2007; Taylor et al. 2008; Kostka et al. 2012). We therefore performed a number of analyses to better understand mechanisms governing rate acceleration of haDHSs. First, to distinguish between relaxation of constraint and true rate acceleration, we applied a novel permutation test (Supplemental Text) and found that 91.8% of haDHSs were evolving faster than their surrounding neutral sequence, suggesting that most haDHSs are not the consequence of relaxed functional constraint. In contrast, it has been estimated that only 55% of HAEs exceed the neutral rate (Kostka et al. 2012). Second, we investigated the contribution of GC-biased gene conversion (GC-BGC) to our data, which influences rate acceleration of HAEs (Pollard et al. 2006a; Galtier and Duret 2007; Duret and Galtier 2009; Kostka et al. 2012), and found that 9.7% (51 haDHSs) show significant evidence of GC-BGC (Supplemental Text; Supplemental Fig. 3a). Finally, we investigated patterns of human–macaque divergence around haDHSs and found that local increases in mutation rate cannot explain rate acceleration in haDHSs, although mutation rate heterogeneity has influenced previous inferences of HAEs (Supplemental Text; Supplemental Fig. 3b; Pollard et al. 2006b; Prabhakar et al. 2006; Bush and Lahn 2008; Lindblad-Toh et al. 2011).

To more directly quantify the proportion of substitutions in haDHSs that can be attributed to positive selection, we used the McDonald-Kreitman framework and compared levels of polymorphism and divergence at haDHSs. Specifically, we used polymorphism data from the 1000 Genomes Project (The 1000 Genomes Project Consortium 2012) and calculated the statistic $\alpha$, an estimate of the proportion of substitutions fixed by adaptive evolution. As a control, we first estimated $\alpha$ in conserved, nonaccelerated DHSs, which as expected was zero (95% CI $-0.02$ to $0.007$) (Fig. 4A; Supplemental Fig. 4a). We estimate that 70.1% (95% CI 65.8%–73.7%) of substitutions can be attributed to positive selection in haDHSs (Fig. 4A), and this number is robust to mutation rate heterogeneity in the presence of complex demographic history (Supplemental Text; Supplemental Fig. 4b). To evaluate the sensitivity of $\alpha$ to GC-BGC, we removed all weak to strong substitutions in haDHSs and repeated the analysis. Although estimates of $\alpha$ decreased for haDHSs subject to GC-BGC, $\alpha$ increased slightly for other haDHSs, and thus the overall estimate remained almost identical (69.9%, 95% CI 64.2%–75.2%) (Fig. 4A). Of the remaining 29.9% of substitutions in haDHSs not accounted for by positive selection, we estimate 9.0% are expected without human-specific rate
acceleration and 20.9% are attributable to additional factors such as relaxation of constraint (Fig. 4B). In support of this hypothesis, we find increased levels of nucleotide diversity in haDHSs and HAEs (Supplemental Text; Supplemental Fig. 5).

**haDHSs are developmental enhancers that exhibit lineage-specific activity**

We performed extensive experimental studies to better understand the functional significance and potential regulatory roles of haDHSs. We found that nine of our haDHSs had previously been tested for in vivo enhancer activity using a transgenic mouse assay (Visel et al. 2007), and we tested nine additional loci. Overall, 13 out of 18 haDHSs were positive for enhancer activity in one or more tissues at the single time point assayed (e11.5) (Supplemental Table 3). These 13 haDHSs were active in a wide range of tissues (Fig. 5A), with the midbrain (n = 7), forebrain (n = 4), branchial arch (n = 4), and limb (n = 4) as the most frequent tissues showing enhancer activity. Patterns of enhancer activity varied from very broad to very tissue specific (Fig. 5A). One interesting example is located on 11p15 and is only active in the branchial arch (Fig. 5A). This haDHS is located in an intron of SOX6, and as we describe below, we find evidence that it contacts the SOX6 promoter. SOX6 is a developmental transcription factor involved in brain, bone, and cartilage development (Lefebvre et al. 1998). Notably, the branchial arch develops into several structures, including the jaw and larynx (Graham 2003), making this haDHS an intriguing candidate that potentially influences traits such as facial morphology and speech.

We also performed luciferase assays to functionally test haDHSs in a more high-throughput manner. Specifically, we experimentally tested 37 haDHSs in SK-N-MC cells (derived from a neuroepithelioma) and 20 haDHSs in IMR90 cells (fetal lung fibroblasts) by assaying for differences in regulatory activity of the human and chimpanzee orthologs using luciferase reporters. We chose SK-N-MC cells as a proxy for other neural cell types, and we chose IMR90 cells because many haDHSs were active in this cell type. Of the 37 pairs of haDHSs tested in SK-N-MC, 14 showed significant enhancer activity (P < 0.05) (Fig. 5B; Supplemental Fig. 6a; Supplemental Table 4). In IMR90, five out of 20 haDHSs showed significant evidence of enhancer function (P < 0.05) (Fig. 5C; Supplemental Fig. 6b; Supplemental Table 4),
Leveraging chromatin contact data to infer putative regulatory targets of haDHSs

Delineating the set of target genes that haDHSs regulate is key to determining their biological consequences and role in human evolution. However, identifying the targets of regulatory sequences poses a significant challenge. Enhancers often regulate distal genes, and in some cases, these may not be the closest genes to the enhancer (van Arensbergen et al. 2014). Chromatin conformation technologies such as Hi-C (Lieberman-Aiden et al. 2009) identify physical contacts between distinct segments of DNA and have been shown to identify long-range interactions between promoters and enhancers (Sanyal et al. 2012). We leveraged high-coverage Hi-C data from human IMR90 fibroblast cells to identify putative regulatory targets of haDHSs using a rigorous statistical method (Ay et al. 2014). We identified 9000 significant contacts for the 524 haDHSs at 40-kb resolution (FDR = 0.01) (Fig. 6A). On average, haDHSs overlap transcription start sites for 3.5 genes, highlighting the potential benefit of using more sophisticated strategies than simply identifying the nearest gene when inferring regulatory targets. We also found that haDHSs contact fewer genes on average than conserved DHSs (permutation \( P = 0.004 \)), suggesting adaptive regulation is more likely to occur when pleiotropic effects are minimized. Furthermore, 119 haDHSs contact one or more transcription factors, and in total 132 distinct transcription factors are contacted by haDHSs. These include SOX6 (see Fig. 5A), RUNX2, and multiple HOX genes, all of which play important roles in development.

We performed a GO enrichment analysis on the set of genes whose transcriptional start sites are contacted by haDHSs. Because haDHSs are a subset of conserved DHSs, we first performed the analysis on conserved DHS contact regions compared with the genomic background. We found that conserved DHS contacts are highly enriched for developmental genes, including those involved in neuron development (Supplemental Table 5), consistent with previous observations about conserved noncoding sequence (Lowe et al. 2011). Next, we tested for GO enrichments in haDHS contact genes using conserved DHS contact genes as the background and found a significant enrichment for developmental terms, including brain and neuron development (corrected \( P < 0.05 \)) (Supplemental Table 5). These results show that haDHSs target genes are enriched for developmentally and neuronally important genes relative to conserved DHSs, which themselves are already highly enriched for these categories.

Three examples of haDHSs and their putative target regions are shown in Figure 6, B through D. All contain transcription factor motifs that are dramatically strengthened or weakened by human-specific substitutions. These haDHSs are likely targets of adaptive evolution as they show no evidence of GC-BGC and are evolving faster than surrounding neutral sequence. Moreover, all three are also active in only a small number of neuronal cell types, such as fetal brain and fetal spinal cord, indicating a potential role in human-specific cognitive phenotypes. Of particular interest is an haDHS on Chromosome 6 that lies in a gene desert 300 kb from POU3F2, a transcription factor that regulates FOXP2 in a human-specific manner (Fig. 6C; Matic et al. 2013). Two of the substitutions in this haDHS strengthen a putative YY1 transcription factor binding site (Fig. 6C), which is known to mediate long-distance DNA interactions (Atchison 2014).

**Discussion**

Advances in DNA sequencing technology have led to a vast catalog of the variation in the genomes and epigenomes across many primates. However, interpreting the evolutionary, functional, and phenotypic significance of these differences and identifying the precise genetic changes that are causally related to human-specific traits remain a formidable challenge. Here, we have leveraged extensive maps of experimentally defined regulatory DNA and comprehensive comparative and population genomics analyses to...
identify and delimit the characteristics of conserved and human accelerated regulatory DNA. In total, we discovered 113,577 DHSs conserved in primates, 524 of which exhibit significant rates of acceleration in the human lineage. We found marked heterogeneity in the distribution of conserved DHSs across cell types (Fig. 2B), with fetal cell types showing the largest amount of constraint. Conversely, DHSs in malignant cell types exhibited the lowest levels of conservation, an observation that may provide insight into cancer biology. For example, chromatin remodeling is disrupted in many cancers (Morin et al. 2010; Jiao et al. 2011). Previous work has shown that DHSs in malignant cell types are more likely to be cell-type specific and have

Figure 5. Experimental assays of enhancer activity in haDHSs. (A) A schematic of the transgenic mouse model is depicted. Rows in the table correspond to each embryonic region, and numbers in parentheses indicate how many of the haDHSs were positive in the region indicated. Columns represent the 13 haDHSs that showed enhancer activity, and gray boxes indicate what tissues the haDHS was active in. Three examples of positive assays are shown above, along with a schematic depicting their location relative to nearby genes. The haDHS tested is shown in red, and other haDHSs in the region are shown in black. (B, C) Results from luciferase assays for haDHSs that showed significant enhancer activity in SK-N-MC and IMR90 cells, respectively. Dotted lines indicate the mean relative expression from the negative controls, and the gray box indicates haDHS human and chimpanzee sequences that showed significantly different activity (P < 0.05). Bars, SE. Asterisks below each plot indicate haDHSs that were active in SK-N-MC or IMR90 (other haDHSs were active in similar cell types, such as fetal brain or NHLF). (D) A schematic of the region surrounding haDHS12, which had the largest difference in enhancer activity. The haDHS is located just upstream of the alternatively spliced gene RNF145. Red substitutions are weak to strong, and all other substitutions are colored in blue. PhyloP scores are also shown across the region. This DHS was partitioned prior to statistical testing into two distinct DHSs. The red portion is human accelerated, and the black portion is not.
levels of nucleotide diversity consistent with neutral evolution (Vernot et al. 2012). Thus, these observations combined with our results that DHSs in malignant cell types have low levels of evolutionary conservation suggest that many malignant DHSs may reflect ectopic chromatin activation.

Our results also provide new insights into human-specific adaptive regulatory evolution. Of the 524 haDHSs that we identified, 454 (87%) are novel and were not detected in previous studies of HAEs (Pollard et al. 2006b; Prabhakar et al. 2006; Bush and Lahn 2008; Lindblad-Toh et al. 2011). The haDHSs that we discovered are significantly less affected by GC-biased gene conversion and relaxation of functional constraint and have a higher proportion of substitutions that are estimated to be due to positive selection compared with previous catalogs of HAEs (Supplemental Fig. 3). We hypothesize these differences are largely the consequence of our study design that synergistically integrated experimentally defined regulatory sequences with phylogenetic conservation, which both focused our analyses to a subset of the genome enriched for functionally important sequence and limited the influence of confounding evolutionary forces. To support this hypothesis, we find that a higher proportion of haDHSs overlap human-specific enhancer marks in the cortex (Reilly et al. 2015) than HAEs ($P = 7.62 \times 10^{-5}$; Fisher’s exact test). Large catalogs of experimentally defined regulatory DNA did not exist when HAEs were initially discovered, and we anticipate that the continued development of functional genomics technology will enable even more refined evolutionary analyses than described here.

To help interpret the functional and potential phenotypic significance of haDHSs, we performed extensive bioinformatics analyses and experimental validations. We found that haDHSs were significantly enriched in noncoding regions; a large proportion of experimentally tested elements showed enhancer activity; and many were active in brain or neural cell types and during fetal development. We also used Hi-C data to inform inferences of putative target genes that are regulated by haDHSs. These analyses revealed that haDHSs contact the transcriptional start sites of 132 transcription factors, suggesting that fine-tuning regulatory

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**Figure 6.** Hi-C chromatin conformation data identify putative regulatory targets of haDHSs. (A) Contacts are shown for all haDHSs, and each row indicates the contacts for one haDHS, which is in the center. Black boxes indicate one 40-kb contact region. The schematic above illustrates how chromatin conformation information gets translated into the Hi-C contact data. Blue dots represent contact regions; the red dot, an haDHS. (B–D) Three example haDHSs are shown with their surrounding genes and a predicted transcription factor binding site that is affected by a human-specific mutation(s). Genes that contact the haDHSs in Hi-C data are highlighted in blue, with arrows pointing to their transcription start sites. Examples B and C depict substitutions that create transcription factor binding sites, while D is a binding site that is predicted to be lost in humans. Human-specific substitutions that go from a weak to a strong base are shown in red, while all other substitutions are shown in blue. Bar plots, FIMO (Grant et al. 2011) log likelihood ratios of motif calls in each species.
networks by tinkering with the sequences that govern the expression of regulatory proteins has been an important target of positive selection during human evolution. A number of transcription factors contacted by haDHSs are strong candidates for influencing hominin- or human-specific traits. For example, RUNX2 has been hypothesized to influence differential bone morphology in humans and Neanderthals (Green et al. 2010), and HOX genes play myriad roles in development. Another intriguing transcription factor contacted by an haDHS is POUSF2, which has recently been shown to regulate FOXP2 in a human-specific manner (Maricic et al. 2013). FOXP2 itself is a transcription factor that has previously been hypothesized to play a role in speech and language in humans (Enard et al. 2002). Our findings suggest that there may be additional levels of human-specific FOXP2 regulation via differential expression of POUSF2 expression. Furthermore, in addition to transcription factors, we identified other genes that are of significant biological interest. For instance, PEX2 is contacted by an haDHS with two substitutions that create a SMAD4 motif (Fig. 6B). Mutations in PEX2 can lead to Zellweger syndrome, characterized by a constellation of features, including impaired brain development and craniofacial abnormalities (Steinberg et al. 2006).

Our study has a number of important limitations. For example, the DHSs we used were ascertained only in human tissues. Although experimentally defined regulatory DNA has been generated in a limited number of nonhuman primates for a limited number of tissues (Shibata et al. 2012; Cotney et al. 2013), a more systematic and comprehensive effort would be of considerable value in understanding the evolution of regulatory sequences. Furthermore, we did not consider additional types of genetic variation, such as structural variation, that may influence human-specific phenotypes (Dennis et al. 2012; Sudmant et al. 2013). Furthermore, although there is evidence that chromatin conformation is relatively stable across cell types (Dixon et al. 2012), it would be of considerable interest to generate Hi-C or related data for a more comprehensive panel of cell types. These data, combined with gene expression profiles from the same tissue types, would provide further insights into the target genes regulated by haDHSs. Finally, the transgenic mouse and luciferase assays that we performed are only a first step in the experimental characterization of these and other elements that potentially contribute to human-specific phenotypes. Because the activity of a regulatory element may be highly cell-type- and developmental time point-specific, and depend on the coordination of additional regulatory elements, more extensive in vivo experiments would be fruitful. Nonetheless, associating particular haDHSs with specific phenotypes is complicated by the fact that the putative causal alleles are fixed in humans and thus refractory to traditional genetic mapping methods. However, if mutations at these sites are not lethal, given the current global population size of humans, such mutations are expected to exist, and their discovery could provide valuable phenotypic insights.

In short, our data provide substantial new insights into sequences that have experienced human-specific adaptive regulatory evolution, narrow the set of genetic changes that may influence uniquely human phenotype, and facilitate more detailed experimental and animal models of the most promising human-specific substitutions.

Ultimately, delineating the suite of genetic changes that have causally influenced human-specific phenotypes will provide insight into the evolutionary and molecular mechanisms that shaped our species evolutionary trajectory.

Methods

DNase I hypersensitivity sites

We used DNase I hypersensitivity peaks previously published as part of the ENCODE (The ENCODE Project Consortium 2012; Maurano et al. 2012) and Roadmap Epigenomics (Bernstein et al. 2010) Projects. A list of cell types is available in Supplemental Table 1. All peaks were called using the hotspot algorithm (John et al. 2011) and represent the 150-bp region of maximal DNase I signal. We merged DHSs across cell types using the BEDOPS package (Neph et al. 2012). Many DHSs were very long after merging (>2000 bp), probably because they consist of distinct regulatory elements located in close succession along the genome. To avoid analyzing distinct, potentially independently evolving regulatory elements as a single unit, we segmented merged DHSs according to the number of cell types each region was active in (Supplemental Text).

Primate alignments

We downloaded the six primate EPO alignment from Ensembl version 70 (Flicek et al. 2014). By use of this, we obtained an alignment for each DHS and the surrounding 50 kb of sequence. We masked all sites that were polymorphic in the 1000 Genomes Project (The 1000 Genomes Project Consortium 2012) integrated phase 1 data (March 2012) at <95% allele frequency, all repeat masked bases (lower case mark up in the EPO alignment), and all sites that were part of a CpG in any species in the alignment. In the surrounding 50 kb, we additionally masked all segmental duplications (UCSC Table Browser), coding exons (UCSC RefSeq genes) padded by 10 bp in order to remove splice sites, promoters (500 bp upstream of transcription start sites), other DHSs, and phastCons Eutherian mammal and primate conserved elements (UCSC phyloP46way). This helped ensure that the 50-kb surrounding region was a more appropriate approximation of the neutral evolutionary model for each DHS. We filtered any DHS in which (1) <90% of the bases remained unmasked in the DHS or (2) <15 kb remained unmasked in any of the six primates in the neutral region. Note, the EPO alignment is based on GRCh37 (hg19), and all subsequent analyses were done using GRCh37 coordinates. Given that we focus on conserved elements, which are by definition located in regions of the genome that are well resolved and alignable, we do not anticipate realigning to GRCh38 would significantly affect our results.

Identifying conserved and accelerated DHSs

DHSs that passed filtering were tested for overall conservation along the primate lineage with software from the PHAST package (Pollard et al. 2010; Hubisz et al. 2011). For each DHS, we first ran phyloFit on the neutral alignment of the surrounding 50 kb with the parameters –nrate 4 –subtree mod SSREV –EM. We used the newick tree provided with the six primate alignment in Ensembl. The resulting file was used as the neutral model while running phyloP; phyloP was run with the parameters –value package (http://github.com/jdstorey/qvalue) for R (R Core Team 2014) were then tested for human acceleration. For this test we used the same neutral model of evolution, this time using the parameters –method LRT –mode CON after removing human sequence from the alignment. DHSs that were conserved at an FDR of 1% as determined with the Q-value package (http://github.com/jdstorey/qvalue) for R (R Core Team 2014) were then tested for human acceleration. For this test we used the same neutral model of evolution, this time using the parameters –method LRT –mode ACC –subtree homo_sapiens. DHSs significant for human acceleration at an FDR of 5% were considered in further analyses. We evaluated the accuracy of the FDR using a sampling approach (Supplemental Text).
To determine the overall rate of evolution in the neutral regions compared with haDHSs, we first concatenated sequence from both sets of regions and then conducted the same set of tests on the regions as a whole. To determine how much faster the human branch in the haDHSs was evolving compared with the expected rate, we multiplied the estimated neutral human branch length by the estimated conservation scale factor and divided the actual haDHS human branch length by this expected number.

Distribution of DHSS across cell types and genomic location

To determine how conserved and accelerated DHSSs were distributed across cell types, we used the bedmap program from the BEDOPS suite (Neph et al. 2012) to map DHSSs from individual cell types onto the set of merged DHSSs. We then calculated the proportion of DHSSs in each cell type that were called as conserved and the proportion of conserved DHSSs that were also called as accelerated (Fig. 2B; Supplemental Fig. 1a–c).

Distribution of DHSSs and haDHSs across the genome was assessed using UCSC Known Gene annotations from the UCSC Genome Browser, downloaded on May 14, 2013. Annotations were filtered to contain only “canonical” transcripts from the knownCanonical table. Promoters were defined as the 500 bp upstream of a transcription start site. To identify physical clusters of haDHSs, we expanded each haDHS by 500 kb on either side and then used the bedmap–count command from the BEDOPS suite (Neph et al. 2012) to count the number of haDHSs and conserved DHSSs within each 1-Mb region.

Other HAEs

We obtained previously identified HAEs (Pollard et al. 2006a,b; Prabhakar et al. 2006; Bush and Lahn 2008; Lindblad-Toh et al. 2011) and assessed overlap using the bedmap program from the BEDOPS package (Neph et al. 2012). When comparing our haDHSs to these other HAEs, we merged all HAEs, again using the BEDOPS program. It was useful for us to compare haDHSs to DHSSs that were conserved but not accelerated. In order to do similar analyses using the HAEs, we merged phastCons elements (UCSC Genome Browser) and considered any element that was >100 bp.

To determine the proportion of overlap between haDHSs and other HAEs, we calculated an empirical null distribution by randomly sampling 524 conserved DHSs 10^4 times and determining overlap with HAEs for each sample.

Population genetics analyses

We downloaded the phase 1 integrated release data from the 1000 Genomes Project (The 1000 Genomes Project Consortium 2012) and filtered sites according to several criteria (Supplemental Text). We calculated θ as described previously (Charlesworth 1994), using the equation 1 – (P_FxP_F), where P = number of polymorphic sites, F = number of human-specific substitutions, S = number of selected sites, and N = number of neutral sites. We considered bases within haDHSs to be putatively selected and bases in the surrounding 4-kb region to be putatively neutral.

Hi-C analyses

We obtained raw paired-end Hi-C libraries for two IMR90 fibroblast cell lines (Dixon et al. 2012). Although Hi-C data were not directly comparable and combined. To accomplish this, we used the lm() function in R (R Core Team 2014) to create a linear model with the ratio of firefly to Renilla for all negative control replicates was a function of plate number. Then the coefficient for each plate was subtracted from all data points for that plate. Enhancer activity was determined using a one-sided f-test, and haDHSs were considered enhancers if either the chimp and/or human allele showed greater luciferase activity than the negative controls. We then line at 40-kb resolution as previously described (Ay et al. 2014). Briefly, we mapped reads to the hg19 (GRCh 37) reference sequence, pairing mapped read ends, filtering duplicates, binning at 40 kb resolution, normalizing raw contact maps (Imakaev et al. 2012), and assigning statistical confidences for each contact bin pair using Fit-Hi-C with a refined null (Ay et al. 2014). We used a significance threshold of q-value <0.01 to determine regions that are contacted by haDHSs containing 40-kb windows. We omitted contacts within the same window and between adjacent windows and only focused on intrachromosomal contacts within 5 Mb of haDHSs. Note that the binning at a coarse resolution and omission of interchromosomal contacts were done to identify only high-confidence contacts with enough sequencing coverage. We used RefSeq gene annotations to obtain a list of transcription start sites that overlap contact regions and used these to perform GO analyses using the WebGestalt server (Wang et al. 2013) with the multiple testing method set to BH and the minimum number of genes per category to 10.

Transgenic mouse assays

Transgenic mouse assays were performed as previously described (Visel et al. 2007). Note, one of the previously tested assays was performed with the mouse ortholog (see Supplemental Table 3). Images of all the mouse assay replicates are available on the VISTA Enhancer Browser (Visel et al. 2007).

Luciferase assays

We considered several factors when selecting which haDHSs to experimentally study. First, because the luciferase assays detect enhancers, we prioritized haDHSs showing evidence of enhancer activity. To this end, we identified a second set of haDHSs that were within 500 bp of an enhancer histone modification (H3K4me1, H3K27ac) signal identified in the same cell type. Histone modifications for this set of haDHSs were downloaded from the UCSC Genome Browser or the Roadmap Epigenomics website. We included only DHSs from the 20 cell types for which histone modification data were available (for additional set of haDHSs and the cell types used, see Supplemental Table 6). There is a column identifying which haDHSs were used in the luciferase assays in Supplemental Tables 2 and 6. Second, we prioritized haDHSs that were active in IMR90, SK-N-MC, or other similar cell types. Both cell types represent time points that are potentially interesting for studying human evolution: SK-N-MC is a brain cell type, and IMR90 is a fetal tissue. Finally, we prioritized haDHSs that showed the greatest evidence for human acceleration.

We used standard techniques for cloning, transfection, and performing luciferase assays. Details are provided in the Supplement. For the luciferase assays, each allele and control had three to eight replicates. The positive control for each plate was cells transfected with the pGL3 control plasmid containing a minimal promoter with strong SV40 enhancer, while the negative control for each plate was cells transfected with the empty pGL3 plasmid with minimal promoter but no additional sequence cloned in.

To increase power to detect enhancer activity, negative control replicates were normalized by plate so that they could be directly comparable and combined. To accomplish this, we used the lm() function in R (R Core Team 2014) to create a linear model with the ratio of firefly to Renilla for all negative control replicates was a function of plate number. Then the coefficient for each plate was subtracted from all data points for that plate. Enhancer activity was determined using a one-sided t-test, and haDHSs were considered enhancers if either the chimp and/or human allele showed greater luciferase activity than the negative controls. We then
tested enhancers for allelic differences with a two-sided t-test between the human and chimp alleles.

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