The characteristics and evolutionary forces acting on regulatory variation in humans remains elusive because of the difficulty in defining functionally important noncoding DNA. Here, we combine genome-scale maps of regulatory DNA marked by DNase I hypersensitive sites (DHSs) from 138 cell and tissue types with whole-genome sequences of 53 geographically diverse individuals in order to better delimit the patterns of regulatory variation in humans. We estimate that individuals likely harbor many more functionally important variants in regulatory DNA compared with protein-coding regions, although they are likely to have, on average, smaller effect sizes. Moreover, we demonstrate that there is significant heterogeneity in the level of functional constraint in regulatory DNA among different cell types. We also find marked variability in functional constraint among transcription factor motifs in regulatory DNA, with sequence motifs for major developmental regulators, such as HOX proteins, exhibiting levels of constraint comparable to protein-coding regions. Finally, we perform a genome-wide scan of recent positive selection and identify hundreds of novel substrates of adaptive regulatory evolution that are enriched for biologically interesting pathways such as melanogenesis and adipocytokine signaling. This data and results provide new insights into patterns of regulatory variation in individuals and populations and demonstrate that a large proportion of functionally important variation lies beyond the exome.

[Supplemental material is available for this article.]

Protein-coding DNA constitutes ~1.5% of the human genome, but ~2.5%–15% is estimated to be functionally constrained (Mouse Genome Sequencing Consortium 2002; Lunter et al. 2006; Asthana et al. 2007; Meader et al. 2010; Ponting and Hardison 2011). Thus, a significant amount of functionally important DNA is located in noncoding regions, and genetic variation in such regions likely makes a significant contribution to phenotypic variation and disease susceptibility among individuals. For example, regulatory variation has been linked to the susceptibility of a wide variety of human diseases, including infectious, autoimmune, psychiatric, neoplastic, and neurodegenerative disorders (for review, see Skelly et al. 2009). In addition, regulatory variation is an important substrate for evolutionary change within and between species (King and Wilson 1975), and a number of examples in humans have been described of positive selection that are due to adaptive evolution of noncoding DNA (Bamshad et al. 2002; Hamblin et al. 2002; Bersaglieri et al. 2004; Tishkoff et al. 2007).

To date, the identification and interpretation of regulatory variation has been challenging because of the difficulty in accurately localizing functional noncoding elements that regulate transcription. Computational approaches for large-scale delineation of regulatory DNA have generally been disappointing. For instance, although sophisticated methods have been developed to identify cis-regulatory motifs, such as transcription factor binding sites and 3′ UTR elements (Hughes et al. 2000; Stormo 2000; Vavouri and Elgar 2005; Xie et al. 2005), it is often unclear how many of the predicted sites are functional. Furthermore, evolutionary-based methods (Siepel et al. 2005; Pollard et al. 2010) are a powerful approach for identifying conserved noncoding DNA that is likely to be functionally important, but only a fraction of such sites encode experimentally verifiable transcriptional control elements, while other data suggest that a large fraction of binding sites for specific regulatory factors is not constrained between species, in part due to lineage-specific use of regulatory elements (Dermitzakis and Clark 2002; The ENCODE Project Consortium 2007, 2012; Blow et al. 2010; Schmidt et al. 2010). Consequently, between-species conservation-based methods likely miss many functional elements.

Although computational and evolutionary-based methods play a critical role in understanding the biology of genomes and interpreting the consequences of putative regulatory variation, experimental methods are the most direct approach for assessing the functional significance of noncoding variation. To this end, large-scale experimental studies of noncoding DNA, harnessing new technologies, such as the ENCODE Project (The ENCODE Project Consortium 2007, 2012) are providing a detailed roadmap to the locations of regulatory DNA in the human genome.

A generic structural feature of animal regulatory DNA is extreme accessibility to nucleases in the context of intact nuclei (Gross and Garrard 1988), and hypersensitivity to the nonspecific endonuclease DNase I has been used for over 30 yr as a probe for regulatory DNA (Galas and Schmitz 1978). The binding of sequence-specific transcriptional regulators in place of canonical nucleosomes creates DNase I hypersensitive sites (DHSs). Nucleotide resolution analysis of DNase I cleavage patterns allows identification of the “footprints” of DNA-bound regulators (Galas and Schmitz 1978). In contrast to ChIP-chip and ChIP-seq, which probe the locations of regulatory sequences for a specific transcription factor, the nonspecificity of DNase I is a powerful feature that allows all DNA–protein interactions to be queried in a single experiment. Large-scale localization of in vivo DNase I cleavages using deep sequencing (Hesselberth et al. 2009) has enabled the
creation of genome-scale maps of diverse functional noncoding
elements marked by DHSs. For example, in the ENCODE Project,
~3 million DHSs have now been mapped across 138 cell types
(Thurman et al. 2012). In addition, genomic DNase I footprinting
of 41 cell diverse cell types has resulted in the localization of 8.4
million DNase I footprints (Neph et al. 2012b).

Here, we describe a comprehensive analysis into patterns of
genetic variation in regulatory DNA marked by DHSs and DNase I
footprints (The ENCODE Project Consortium 2012). By analyzing
whole-genome sequence data, we are able to directly compare
characteristics of regulatory and protein-coding variation and find
that individuals harbor considerably more regulatory compared to
protein-coding variants. Moreover, we demonstrate that signifi-
cant heterogeneity of functional constraint exists across regulatory
DNA between cell types and that regulatory DNA present in mul-
tiple broad categories of cell types is significantly more con-
strained. Finally, we quantify patterns of population structure in
regulatory DNA and identify several hundred loci that contain
signatures of local adaptation. In summary, these analyses represent
the most comprehensive assessment of human regulatory variation
described to date and have important implications for personal
genomics, disease mapping studies, and human evolution.

Results and Discussion

Overview of DNase I and whole-genome sequence data

A schematic illustration of the classes of data used in our analyses is
shown in Figure 1A. Within DHSs, DNase I “peaks” correspond to
−150-bp regions of maximum hypersensitivity (Fig. 1A; see The
ENCODE Project Consortium 2012). Embedded within peaks, are
much smaller 6- to 20-bp DNase I footprints, which identify
regions bound by regulatory factors (Fig. 1A). We also obtained
publicly available whole-genome sequence data for 53 unrelated
individuals that encompass five geographically diverse popula-
tions (Fig. 1B) from Complete Genomics. The average sequencing
depth per individual was ~40×. Variants were filtered for devi-
ations from Hardy-Weinberg equilibrium, and partial genotype
calls were set to missing data (see Methods). The high-coverage
whole-genome sequence data are ideal for population genetics
analyses as they are free from the confounding affects of ascer-
tainment bias present in genotypes obtained from SNP chips
(Tennessen et al. 2011).

Pervasive regulatory variation across the human genome

Across all cell types, over 2.9 million DNase I peaks and 8.4 million
DNase I footprints were identified across sampled cell types that
collectively span 577 Mb and 156 Mb of sequence, respectively
(18.7% and 5.1% of the genome for peaks and footprints, respec-
tively). By use of the whole-genome sequence data, we observed
3.85 million, 1.01 million, and 0.15 million variants in DNase I
peaks, DNase I footprints, and the exome, respectively. The large
number of variants in peaks and footprints relative to exomes is
a function of the total amount of sequence they encompass. For
example, the number of variants per kilobase in peaks, footprints,
and the exome is 6.7, 6.5, and 4.2, respectively.

To compare the number of putatively functional variants
across peaks, footprints, and the exome we obtained GERP scores
for each variant, which is a measure of evolutionary constraint
with positive values indicating greater conservation (Cooper et al.
2005). Peaks and footprints not only have an overall larger number
of variants relative to exomes but also manifest more high GERP
variants compared with protein-coding regions (Fig. 2A), although
the differences between categories becomes less dramatic. For ex-
ample, at a threshold of GERP ≥ 3 (Cooper et al. 2005) 146,570,
61,933, and 36,935 variants are observed in peaks, footprints,
and the exome, respectively. It is interesting to note that protein-coding
DNA contains proportionally more putatively functional variation
compared with noncoding DNA (i.e., 24.6%, 6.1%, and 3.8%
of variants have a GERP ≥ 3 for exomes, footprints, and peaks,
respectively), but the absolute number of functional variants in

Figure 1. Overview of data used in the analyses. (A) Schematic of the DNase I data. Binding of regulatory proteins to DNA (blue rectangle) results in
nucleosome (open circles) displacement and local chromatin remodeling, and these regions are susceptible to cleavage with the endonuclease DNase I.
High-throughput sequencing of libraries made from digested nuclei reveals DNase I hypersensitive sites, detectable by increased depth of coverage. Peaks
are defined as 150-bp windows centered on the area of maximum cleavage (The ENCODE Project Consortium 2012). Within hypersensitive sites,
footprints of regulatory factor binding are observed as decreased cleavage. (B) Unrooted neighbor-joining tree of the 53 unrelated individuals colored by
population. Abbreviations are described in Supplemental Table 2.
noncoding regions is larger because of the greater amount of genomic sequence they encompass. Thus, regulatory variation is pervasive across the human genome, and a substantial proportion of functional variation exists in noncoding DNA.

Next, we investigated the distribution of putative regulatory and protein-coding variation across individuals. As expected, the average number of variants (±SD) per individual in peaks and footprints is dramatically higher than that found in the exome (741k ± 72k in peaks, 192k ± 18k in footprints, and 24.4k ± 2.2k in the exome) (Fig. 2B). A more interesting comparison, however, is the number of putatively functional regulatory and protein variants per individual. Therefore, we also determined the number of variants per individual with a GERP ≥ 3 in peaks, footprints, and the exome (Fig. 2C). On average, individuals contain 24.2k ± 2.3k, 10.1k ± 0.92k, and 4.7k ± 0.40k high GERP variants in peaks, footprints, and the exome, respectively (Fig. 2C). Although evolutionary constraint is not a perfect proxy for function, these results suggest that individuals possess more regulatory versus protein-coding variants. Assuming the probability that a variant is functional is the same between coding and noncoding DNA, this assumption, however, is dubious (McVicker et al. 2009), and more definitive inferences on the proportion of functional variants in noncoding versus coding DNA will ultimately require further experimental data. In addition, it is interesting to note that, as expected, the average number of variants per individual in peaks and footprints is significantly higher for individuals of African ancestry compared to non-Africans (859k vs. 710k in Africans and non-Africans, respectively; $P < 9.95 \times 10^{-10}$) (Fig. 2B).

Patterns of nucleotide diversity in regulatory DNA sequence motifs

The unique scope of the data sets analyzed here allows us for the first time to systematically investigate genomic patterns of variation in DNA sequence motifs. To this end, we scanned DNase I footprints for 732 known motifs (see Methods), and for each motif, we calculated nucleotide diversity, $\pi$, averaged across all instances of the motif in these regions. To facilitate interpretation of motif diversity, we also calculated $\pi$ for fourfold synonymous sites, a proxy for neutrally evolving DNA, and protein-coding sequences. As shown in Figure 3A, average diversity varies by over sevenfold across known regulatory motifs, ranging from $2.67 \times 10^{-4}$ to $2.0 \times 10^{-3}$. Approximately 60% of motifs have average diversities significantly lower than fourfold synonymous sites (Fig. 3A), indicative of purifying selection.

Figure 3A also highlights motif diversity for several important classes of transcriptional regulators. For example, HOX-, POU-, and FOX-domain factors are heavily enriched in developmental regulators and controllers of cellular differentiation. Motifs for transcription factors belonging to these classes are markedly shifted toward lower diversity, and motifs for several individual factors exhibit levels of diversity that are reduced beyond that of protein-coding sequences (Fig. 3A). In contrast, diversity in motifs for tandem zinc finger transcription factors, which comprise the largest and most diverse class of human transcription factors, is distributed relatively evenly across the diversity spectrum (Fig. 3A). Members of this group include core regulatory factors such as CTCF and YY1, developmental regulators such as PRDM1 and ZIC3, and numerous chromatin repressors such as RREB1, REST, and the KRAB-ZNF family of proteins. Because many of the canonical motifs for these factors contain one or more CG dinucleotides, we hypothesized that the increased average diversity for these factors might be a consequence of higher mutation rates at CpG sites. To explore this hypothesis, we identified factors for which >50% of the motif instances in regulatory DNA contained CpGs, which revealed that the ubiquitous presence of CpG sites is a common characteristic of motifs with high levels of diversity (Fig. 3A).

To more systematically control for mutation rate heterogeneity, we also calculated $\pi$ normalized for divergence (see Methods) for each motif. As shown in Figure 3B, normalized diversity has the
Heterogeneity of functional constraint across cell types

We next investigated differences in normalized diversity between “core” DHS and DHS found in only one category of cell types. To this end, all of the cell types can be grouped into one of three categories: normal/primary, iPS/ES, and malignant. To minimize potential contributions from experimental noise, we focused on a subset of 92 cell types with high-quality DNase I data in which >40% all sequence tags map within DHSs (equivalent to average signal-to-noise of \(~100:1\) (Thurman et al. 2012) and calculated normalized $\pi$ in DNase I peaks that are shared and unique to each cell type category (Fig. 4B). Eight percent of peaks are found in all three categories, whereas 6.4%, 31.1%, and 28.2% of peaks are unique to iPS/ES, malignant, and normal/primary cell types, respectively (Fig. 4B). Overall, there is significant variation ($P < 10^{-16}$) in normalized diversity among peaks shared between cell type categories versus those found in a single category (Fig. 4B). In particular, DNase I peaks shared by two or three categories of cell types exhibit the lowest levels of normalized diversity (Fig. 4B), consistent with stronger selective constraint. Conversely, peaks found in only one cell category have significantly higher normalized diversity ($P < 10^{-1}$) (Fig. 4B). These results suggest that the “core” set of DHSs, present in more than one cell type category, is subject to stronger purifying selection because they are necessary for proper transcriptional programs in multiple cell types.
Evidence for ectopic activation of DHSs in malignant cell types

Many cancers are characterized by disruptions in chromatin maintenance pathways (Wang et al. 2007; Morin et al. 2010; Jiao et al. 2011). Additionally, many immortalized cells express different complements or ratios of transcriptional factors than are found in normal differentiated cells (Zaidi et al. 2007). These observations suggest that immortalized and malignant cell lines may experience increased “ectopic” activation of DHSs. To explore the potential ectopic activation of DHSs in malignant and immortalized cell types further, we calculated the proportion of DNase I peaks that are present in only one cell line, as noncanonical chromatin remodeling would be expected to result in an excess of cell type–restricted DHSs. Again, we used the same 92 cell types as described above.

We found that 54% of peaks specific to normal/primary cells are present in a single cell type. In contrast, 81% of malignant-specific peaks and 86% of iPS/ES-specific peaks are present in a single cell line. However, these percentages are not directly comparable, because of sample size differences between categories (n = 58, 29, five for normal/primary, malignant, and iPS/ES, respectively). When we correct for the number of cell types per category (see Methods), we find that iPS/ES cells are not enriched for singleton DHSs compared to normal/primary cells (P = 0.236), whereas malignant cell types are significantly enriched (P < 10^-4) for singleton DHSs compared to normal/primary cells (Fig. 5). These data raise the intriguing possibility that the DHSs found in malignant cells, though not increased significantly in number (data not shown), are enriched in elements resulting from ectopic cooperative transcription factor binding within neutrally evolving sequences.

Signatures of positive selection

A large number of genome-wide scans for recent positive selection have been performed in humans (for review, see Akey 2009). Typically, these studies focus only on patterns of DNA sequence variation and are not informed by functional genomics data, although genome-wide analyses have been pursued on computa-
tionally predicted motifs. The large compendium of experimentally characterized regulatory regions provides a unique data set to interrogate for signatures of recent positive selection.

To this end, we performed a population genomics analysis to identify DNase I peaks that contain variants with large allele frequency differences between populations relative to the genomewide, which is a signature of geographically restricted selection (Akey et al. 2002; Akey et al. 2004). Specifically, we calculated locus-specific branch lengths (LSBLs) (Shriver et al. 2004) for variants in DNase I peaks in Africans, Asians, and Europeans. LSBL is a function of pairwise $F_{ST}$ between populations (see Methods) and helps isolate the direction of allele frequency change (Shriver et al. 2004). To reduce the stochasticity inherent in summary statistics of population differentiation, we averaged LSBL across all variants in a peak. We excluded X-linked variants from our analysis due to its different effective population size.

The genome-wide distributions of population structure in DNase I peaks in the African, Asian, and European populations are shown in Figure 6A. We pursued two distinct approaches to interpret these data. First, to obtain general insights into the characteristics of DNase I peaks that exhibit large allele frequency differences between populations, we focused on peaks in the 1% tail of the empirical distribution of LSBLs in each population (Fig. 6A). Next, we identified all genes within 50 kb of these peaks ($n = 3372, 3224, \text{and } 3099$ such genes in Africans, Asians, and Europeans, respectively) and tested for enrichment of KEGG pathways. As shown in Table 1, this set of genes is significantly enriched for 15 KEGG pathways, seven of which are shared between two or more populations (including pathways related to cancer, axon guidance, and WNT signaling). Interestingly, the most significantly enriched pathway in Europeans is melanogenesis (Table 1), suggesting that in addition to protein-coding variants (Lamason et al. 2005), regulatory polymorphisms influencing pigmentation phenotypes have also been a target of recent positive selection. Moreover, our African sample is significantly enriched for chemokine and adipocytokine signaling pathways (Table 1), which is particularly interesting given the known differences in prevalence of insulin resistance and type 2 diabetes in individuals of African ancestry (Reimann et al. 2007).

**Figure 6.** Genome-wide distribution of population structure in regulatory DNA. (A) Genome-wide distribution of locus-specific branch lengths (LSBLs) for Africans, Asians, and Europeans, respectively. Note that the valley of uniform LSBL on chromosome 17 in Europeans corresponds to the MAPT region that is segregating a large chromosomal inversion (Zody et al. 2008). (B) Distribution of the proportion of highly differentiated DNase I peaks found for different categories of cell types. (SC) Stem cells (iPS/ES); (I) immortalized; (M) malignant; (N) normal/primary cell types. (C) Distribution of African LSBL across intron 1 of VDR. (D) Distribution of European LSBL across intron 4 of FTO. In panels C and D, peaks are shown as red rectangles and exons as black rectangles.
We also investigated the distribution of DNase I peaks that exhibit unusually large levels of population structure across cell types. To this end, we classified the 138 types into normal, immortalized, malignant, and pluripotent (iPS/iES) categories. The proportion of DNase I peaks that are in the 1% tail of the empirical distribution of LSLBs is significantly different across cell type categories (Kruskal-Wallis test, $P = 3.2 \times 10^{-15}$). Primary/normal cell lines had the highest proportion of differentiated peaks, whereas iPS/iES cell lines had the lowest proportion of differentiated peaks (Fig. 6B). The higher proportion of differentiated DNase I peaks in primary/normal cell lines is driven by a wide variety of cell types, including astrocytes (spinal cord [HA-sp], cerebellar [HA-c], and cortical [HA-h]), renal glomerular endothelial cells (HRGEC), and cardiac fibroblasts (HC.Faa). Although these results are intriguing and offer preliminary insights into the types of tissues that contribute to fitness differences among individuals, more definitive inferences will require an even broader sampling of cell types.

Second, to develop a more refined list of putative targets of recent adaptive evolution, we focused on the most differentiated 1% of DHSs that also contain one or more highly differentiated variants with a GERP $\geq 3$. In total, 323, 349, and 313 DHSs meet these criteria in Africans, Asians, and Europeans, respectively. We identified genes located within 50 kb of each of these peaks and identified 187, 174, and 179 genes in Africans, Asians, and Europeans, respectively (Supplemental Text 1). Notably, included in this set of peaks is the well-documented promoter variant in DARC that results in malaria resistance in African populations (Hamblin et al. 2002), which demonstrates the potential power of this data set to fine-scale map signatures of selection and identify selected alleles. Moreover, 61, 40, and 51 of these candidate selection genes in Africans, Asians, and Europeans, respectively, overlap previously reported signatures of selection collected by Akey (2009), which is significantly more than expected by chance ($P < 10^{-6}$ for all populations).

Thus, these observations suggest that the loci identified here are enriched for targets of recent positive selection. Particularly interesting examples of novel targets of selection include the vitamin D receptor (VDR) in Africans and the fat mass and obesity associated gene (FTO) in Europeans (Fig. 6C,D). The list of all candidate selection genes located within 50 kb of highly differentiated peaks is provided in Supplemental Text 1, which provides a powerful framework for more detailed analyses into recent adaptation of noncoding DNA in humans.

### Conclusions

By synergistically integrating whole-genome sequences with genome-wide DNase I data, we provide new insights into the distribution and characteristics of human cis-regulatory variation in individuals and populations. Our results demonstrate that regulatory variation is pervasive throughout the genome, on average mildly deleterious, and individuals likely harbor more functionally important variants in noncoding compared with protein-coding DNA. The latter observation is important for disease mapping studies and suggests that a substantial proportion of disease alleles exist beyond the exome (Bamshad et al. 2011). Our results also suggest that ectopic activation of noncanonical cis-regulatory sequences contributes to the aberrant transcriptional changes that are observed in many cancers. Finally, we describe a large compendium of DHSs that exhibit unusually large levels of population structure, consistent with the action of geographically restricted selection. Genes adjacent to these highly differentiated regulatory sequences are enriched for a number of biologically interesting categories, such as signaling and disease related pathways. Although considerably more work is needed to elucidate the evolutionary history of these loci, they provide an important starting point for understanding how recent adaptive evolution has influenced regulatory networks.

### Methods

#### DNase I data

We obtained DNase I peaks, footprints, and predicted motif locations from the ENCODE Project (The ENCODE Project Consortium 2012, http://genome.ucsc.edu/ENCODE/downloads.html). Peaks and footprints were empirically thresholded at a 1% false-discovery rate. For aggregate analyses over DHS across cell types, peak or footprint locations were merged across cell types using BEDOPS (http://code.google.com/p/bedops/) (Neph et al. 2012a). More information about the cell lines can be found in Supplemental Table 1 and http://genome.ucsc.edu/ENCODE/cellTypes.html.

Protein binding motif locations were generated genome-wide with FIMO motif scanning software (Bailey et al. 2009), version 4.6.1, using a $P$-value threshold of $1 \times 10^{-5}$ threshold. Motif models were obtained from TRANSFAC (Matys et al. 2002), version

### Table 1. Enriched KEGG pathways for genes within 50 kb of highly differentiated DNase I peaks

<table>
<thead>
<tr>
<th>Population</th>
<th>KEGG pathway (Identification)</th>
<th>No. of genes</th>
<th>$P$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>European</td>
<td>Melanogenesis (04916)</td>
<td>31</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>Arrhythmogenic right ventricular cardiomyopathy (05412)</td>
<td>28</td>
<td>0.0006</td>
</tr>
<tr>
<td></td>
<td>ECM-receptor interaction (04512)</td>
<td>30</td>
<td>0.0007</td>
</tr>
<tr>
<td></td>
<td>Pathways in cancer (05200)</td>
<td>80</td>
<td>0.0033</td>
</tr>
<tr>
<td></td>
<td>Dilated cardiomyopathy (05414)</td>
<td>29</td>
<td>0.0052</td>
</tr>
<tr>
<td></td>
<td>Hypertrophic cardiomyopathy (05410)</td>
<td>27</td>
<td>0.0066</td>
</tr>
<tr>
<td></td>
<td>Axon guidance (04360)</td>
<td>37</td>
<td>0.0075</td>
</tr>
<tr>
<td></td>
<td>Focal adhesion (04510)</td>
<td>51</td>
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</tr>
<tr>
<td></td>
<td>Wnt signaling pathway (04310)</td>
<td>41</td>
<td>0.0087</td>
</tr>
<tr>
<td></td>
<td>Calcium signaling pathway (04020)</td>
<td>46</td>
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</tr>
<tr>
<td></td>
<td>Vascular smooth muscle contraction (04270)</td>
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<td>0.0008</td>
</tr>
<tr>
<td></td>
<td>Chemokine signaling pathway (04062)</td>
<td>57</td>
<td>0.0008</td>
</tr>
<tr>
<td></td>
<td>Adherins junction (04520)</td>
<td>29</td>
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<td>Pathways in cancer (05200)</td>
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<td>Wnt signaling pathway (04310)</td>
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<tr>
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<td>83</td>
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<td>Hypertrophic cardiomyopathy (05410)</td>
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</tr>
<tr>
<td></td>
<td>Dilated cardiomyopathy (05414)</td>
<td>30</td>
<td>0.0059</td>
</tr>
</tbody>
</table>

$P$-values shown are adjusted for multiple testing using FDR (see Methods). Pathways in bold denote those shared between two or more populations.
2011.1. For these analyses, all motifs were intersected with footprint data using BEDOPS.

Sequence data

We obtained whole-genome sequence data from 69 individuals that were sequenced to high coverage by Complete Genomics (http://www.completegenomics.com/sequence-data/download-data/). Among these 69 individuals, 54 are reported to be unrelated. To verify that these 54 individuals are unrelated, we performed relationship inference with KING (Manichaikul et al. 2010). Two Maasai individuals (NA21732 and NA21737) who were not reported as being related were found to be either siblings or parent–child. We removed NA21737 from further analyses as this individual had more missing data than NA21732. Thus, our final data set consists of 53 individuals from five populations (Supplemental Table 2). Genotype data were filtered to remove partial genotypes (i.e., where one allele is called and the other is reported as missing), by coverage (>20% of individuals must have calls), and by extreme departures from Hardy-Weinberg Equilibrium ($P < 10^{-8}$, which corresponds to all individuals being heterozygous and therefore most likely a paralogous sequence variant). We defined fourfold degenerate sites using NCBI-called reading frames. We used the NimbleGen SeqCap EZ Exome, version 2.0, definition, downloaded from the NimbleGen website (http://www.nimblegen.com/products/seqcap/ez/v2/). Repeats were defined by RepeatMasker regions, obtained from the UCSC Genome Browser. A dinucleotide is conservatively called as CpG for NpG and CpN dinucleotides (where N = A,C,T, or G) in any of our 53 genomes, chimpanzee, orangutan, rhesus macaque.

Statistical analyses

We calculated nucleotide diversity as: $\pi = \frac{1}{2N} \sum_{i=1}^{N} (2p_i / (1 - p_i^2))$, where $n$ is the number of chromosomes and $p_i$ is the frequency of the major allele for the $i$th segregating site, $S$. To obtain a per nucleotide estimate of $\pi$, we divided by the total number of bases considered for a particular analysis. Normalized diversity was calculated by dividing the per nucleotide $\pi$ estimate by the estimated neutral mutation rate. For exonic sequence, we used the mutation rate calculated at fourfold synonymous sites, as this sequence is less likely to be influenced by selection compared to all synonymous sites. For DHSs, we expanded each region (peak or motif) by 1500 bp on either side and removed putatively selected sequence (footprints, exome, and peaks padded by 250 bp) from this region. All normalized $\pi$ values were then multiplied by $2 \times 10^{-8}$ to bring them into the range of non-normalized $\pi$ values. Repeats were removed in all analyses. For $\pi$ in DHSs (normalized and non-normalized), exonic sequence was also removed. CpGs were removed in all normalized $\pi$ calculations as described above.

To evaluate the number of singleton peaks in malignant and iP/iES cells relative to normal/primary cell lines, we performed a resampling procedure. Specifically, we randomly selected 1000 samples of five and 29 cell types from the 58 normal cell lines and, for each sample, calculated the proportion of singleton normal-specific DNase I peaks. Singleton peaks must occur in category-specific DHS; therefore, we calculate the percentage of category-specific DHS that is also singleton. This procedure generates an empirical distribution for the proportion of singleton peaks expected in a category with five or 29 cell types.

We calculated LSBLs as previously described (Shriver et al. 2009). In brief, pairwise $F_{ST}$ between Africans ($n = 16$), Europeans ($n = 13$), and Asians ($n = 8$) was calculated as $1 - H_2/H_1$, where $H_2$ and $H_1$ denote average subpopulation heterozygosity and total heterozygosity, respectively (Hartl and Clark 1997). These pairwise estimates of $F_{ST}$ were then used to calculate LSBL for each population. For example, denote the $F_{ST}$ between Africans and Europeans, Africans and Asians, and Europeans and Asians as $d_{AB}$, $d_{AC}$, $d_{AB}$, respectively. The LSBL for Africans is $d_{AB} + d_{AC} - d_{AB}/2$. LSBLs were calculated for all variants in peaks with 100% coverage (all individuals fully called), excluding the exome and repeats.

To test for enrichment of candidate selection genes in KEGG pathways, we used WebGestalt (Duncan et al. 2010; http://bioinfo.vanderbilt.edu/webgestalt). In these analyses, we used a background list of genes by identifying the closest gene to each DNase I peak, which recapitulates how genes were associated with the highly differentiated peaks. We used the false discovery rate (FDR) method of Benjamini and Hochberg (1995) to address multiple testing.

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