Fine-mapping the Favored Mutation in a Positive Selective Sweep

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Abstract

Methods that scan population genomics data to identify signatures of selective sweep have been actively developed, but mostly do not identify the specific mutation favored by the selective sweep. We present a method, iSAFE that uses population genetics signals and a boosting approach to pinpoint the favored mutation even when the signature of selection extends to 5Mbp. iSAFE was tested extensively on simulated data and 22 known sweeps in human populations using the 1000 genome project data with some evidence for the favored mutation. iSAFE ranked the candidate mutation among the top 13 (out of ~21,000 variants) in 14 of the 22 loci, did not show a strong signal in 3. We identified previously unreported mutations as being favored in the remaining 5 regions. For these pigmentation related genes, iSAFE identified identical selected mutations in multiple non-African populations suggesting an out-of-Africa onset of selection.

Introduction

Genetic data from diverse human populations have revealed a multitude of genomic regions believed to be evolving under positive selection. For the most part, these regions follow a regime where a single, favored, mutation increases in frequency in response to a selection constraint. The favored mutation either exists as standing variation at the onset of selection pressure, or arises de novo, after the onset. Neutral mutations on the same lineage as the favored mutation, hitchhike (are co-inherited) with the favored mutation, and increase in frequency, leading to a loss of genetic diversity.

Methods for detecting genomic regions under selection from population genetic data exploit a variety of genomic signatures. Allele frequency based methods analyze the distortion in the site frequency spectrum; Linkage Disequilibrium (LD) based methods use extended homozygosity in haplotypes; population differentiation based methods use difference in allele frequency between populations; and finally, composite methods combine multiple test scores to improve the resolution. Recently, a lack of rare (singleton) mutations has been used to detect very recent selection. The signature of the selective sweep can be captured even when standing variation or multiple de novo mutations create a ‘soft’ sweep of distinct haplotypes carrying the favored mutation. Together
with the advent of deep sequencing, these methods have identified multiple regions under selection in humans and other organisms, and provide a window into genetic adaptation and evolution.

In contrast, little work has been done to identify the favored mutation in a selective sweep. Grossman et al. note that different selection signals identify overlapping but different regions, and a composite of multiple signals (CMS) can localize the site of the favored mutation. An alternative strategy is to use functional information to annotate SNPs and rank them in order of their functional relevance. However, the signal of selection is often spread over a large region, up to 1–2 Mbp on either side, and the high LD makes it difficult to pinpoint the favored mutation. Here, we propose a method, iSAFE (integrated Selection of Allele Favored by Evolution), that exploits coalescent based signals in the ‘shoulders’ of the selective sweep to rank all mutations within a 5Mbp around a region under selection. iSAFE requires that the broad region under selection is identified using existing methods, but does not depend on knowledge of the specific phenotype under selection, and does not rely on functional annotations of mutations.

Results

iSAFE considers only biallelic sites. It takes as input a binary SNP matrix with each row corresponding to a haplotype \( h \), each column to a site \( e \). Entries in the matrix correspond to the allelic state, with 0 denoting the ancestral allele, and 1 denoting the derived allele. A haplotype ‘contains/carry a mutation \( e \)’ if it has the derived allele at site \( e \). Recently, we devised the Haplotype Allele Frequency (HAF) score to capture the dynamics of a selective sweep. The HAF score for a haplotype \( h \) (HAF(\( h \))) is the sum of the derived allele counts of mutations in \( h \) (Fig. 1A and online methods). It has been shown when \( h \) is a carrier of the favored allele, HAF(\( h \)) increases with the frequency of the favored mutation, in contrast to HAF scores of non-carriers, and this can be used to separate carrier haplotypes from non-carriers without knowing the favored mutation.

Denote two haplotypes as ‘distinct’ if they have different HAF-scores. For any mutation \( e \), let \( f_e \) denote the mutation frequency, or the fraction of haplotypes carrying the mutation. Let \( \kappa(e) \) (Fig. 1B) denote the fraction of distinct haplotypes that carry mutation \( e \).

\[
\kappa(e) = \frac{\# \text{ of distinct haplotypes carrying mutation } e}{\# \text{ of distinct haplotypes in sample}}.
\]  

(1)

Similarly, let \( \phi(e) \) denote a normalization of the sum of HAF-scores of all haplotypes carrying the mutation \( e \).

\[
\phi(e) = \frac{\text{sum of HAF-scores of haplotypes carrying mutation } e}{\text{sum of HAF-scores of all haplotypes}}.
\]  

(2)

We observe empirically that in a region evolving according to a neutral Wright-Fisher model, \( \kappa(e) \) and \( \phi(e) \) are both estimators of \( f_e \), with variance \( f_e(1-f_e) \) (Fig. S1). Based on this observation, we define the SAFE-score of mutation \( e \) as

\[
\text{SAFE}(e) = \frac{\phi(e) - \kappa(e)}{\sqrt{f_e(1-f_e)}}.
\]  

(3)
Empirically, SAFE($e$) behaves like a standard normal random variable under neutrality (Fig. S2), and it can be used to test departure from neutrality. However, its real power appears during positive selection, when SAFE-scores change in a dramatic, but predictable, manner (Fig. 1B,C,D,E). Assuming a no recombination scenario, label mutations as ‘non-carrier’ if they are carried only by haplotypes not carrying the favored allele. The remaining mutations can be labeled as ‘ancestral’, if they arise before the favored mutation, or ‘descendant’, if they arise after (Fig. 1C). Representing each mutation as a point in a 2-dimensional plot of $\phi,\kappa$ values, these classes are clustered differentially (Fig. 1D,E). The selective sweep reduces the number of distinct haplotypes carrying the favored mutation (lower $\kappa$), leaving non-carrier mutations with increased fraction of distinct haplotypes (higher $\kappa$). On the other hand, increased HAF-scores in carrier haplotypes reduces the proportion of total HAF-score contributed by non-carrier haplotypes (lower $\phi$). In contrast, the favored mutation has high positive value of $\phi - \kappa$ due to high HAF-scores for carriers (higher $\phi$), and the reduced number of distinct haplotypes among its descendants (lower $\kappa$). As we go up to ancestral mutations, the number of non-carrier haplotype descendants increase, and $\kappa$ grows faster than $\phi$. As we go down to descendant mutations, there is a reduction in the already small number of distinct haplotypes. However, $\phi$ decreases sharply, reducing $\phi - \kappa$ (see Fig. 1B,C,E). Thus, we expect that the mutation with the highest SAFE-score is a strong candidate for the favored mutation.

We performed extensive simulations\textsuperscript{7} to test SAFE on samples evolving neutrally and under positive selection. We varied one parameter in each run (online methods), including window size ($L = 50$kbp), number of individual haplotypes ($n = 200$) chosen from a larger effective population size ($N = 20K$), scaled selection coefficient ($Ns = 500$), initial and final favored mutation frequencies ($\nu_0 = 1/N$, and $\nu$). No available tool explicitly identifies the favored mutation in a selective sweep. However, the integrated Haplotype Score (iHS) scores each variant according to its likelihood of being under selection in an ongoing sweep, with the goal of detecting regions under ongoing selection\textsuperscript{8}. To provide a baseline for comparison, we compared SAFE against iHS, using iHS scores to rank mutations.

While standing variation, $\nu_0 > 1/N$, generally weakens the selection signal, the performance of SAFE remains relatively robust to variation in $\nu_0$ (Figure 1F). The median SAFE rank of the favored allele is at most 3 out of $\sim250$ variants in all cases except when $\nu_0 \geq 1000/N$. Similarly, the performance is robust to selection pressure, with only a slight degradation at weak selection ($Ns = 50$) (Fig. S4) where the median rank goes from 4 (1.6%-ile) to 9 (3.5%-ile). For $Ns \geq 200$ the median rank is at most 2. As expected, the performance improves with increasing sample size (Fig. S5). We also tested SAFE on a model of European demography and found no considerable loss of performance (Fig. S6). These tests used $L = 50$kbp, chosen so as to minimize the effects of recombination. In testing SAFE on larger regions, we found that while the median rank of the favored mutation increases with increasing window size, the percentile rank improves up to 80kbp and then degrades to $3\%-ile$ around 1Mbp (Fig. 2A, and S7). The deterioration for larger windows is likely due to most haplotypes becoming unique, and $\kappa$ losing its utility in pinpointing the favored
The selective sweep signal can extend to large, linked regions, as far as 1Mbp on either side of the favored allele. These ‘soft-shoulders’ of selective sweeps are helpful in identifying the region under selection, but make it harder to pinpoint the favored mutation. We further refined our method to exploit the signal from soft shoulders.

In analyzing large genomic regions, we considered a set of 50% overlapping windows of fixed size (300 SNPs). For each window, we applied SAFE and chose the mutation with the highest SAFE-score. Let \( S_1 \) denote the set of selected mutations. The favored mutation is the true classifier for carriers (with high haplotype homozygosity/high haplotype counts per unique haplotype) and non-carriers (with low haplotype homozygosity/low haplotype counts per unique haplotype) in the vicinity of the favored mutation. Therefore, the SAFE-score can be considered as a measure of goodness of classification (Fig. 2D). Mutations in \( S_1 \) are likely to contain either the favored mutation itself or mutations linked to it. Moreover, if the true favored (or tightly linked) mutation in window \( w \) is inserted artificially into a different window \( w' \), it will have a high SAFE-score only when the genealogies of \( w, w' \) are identical or very similar, but not otherwise. Other mutations are not expected to have a high SAFE-score when added to any window other than their own (Fig. 2D). We use this insight combined with the idea of boosting with weak classifiers to develop a method for finding the mutation that can best separate the haplotypes into carriers with higher and non-carriers with lower haplotype homozygosity in the region. Let \( \Psi_{e,w} \) denote the larger of the SAFE-score of \( e \), when \( e \) is ‘inserted’ into window \( w \), or 0 (Fig. 2C). Define the weight of a window \( w \) as

\[
\alpha(w) = \frac{\sum_{e \in S_1} \Psi_{e,w}}{\sum_{w' \in W} \sum_{e \in S_1} \Psi_{e,w'}}.
\]

(4)

Windows that contain the favored mutation and those surrounding it are expected to have high \( \alpha \) values. We defined the iSAFE-score for all mutations \( e \) (including those not in \( S_1 \)) as:

\[
iSAFE(e) = \sum_{w \in W} \Psi_{e,w} \cdot \alpha(w).
\]

(5)

We tested the power of iSAFE to identify the favored mutation in varying window sizes and saw little or no loss of performance as the window size was increased from 250kbp all the way to 5Mbp (Fig. 2B). The median rank remains between 3 and 5 up to 5Mbp, and its performance remains robust to a large range of parameter choices including both hard and soft sweep scenarios, selection pressure and favored mutation locations (Fig. S8-S13).

While iSAFE-scores do not have a direct probabilistic interpretation, they are normalized and can be compared across samples. We found distinct differences in performance after a score threshold of 0.1. The median rank of the favored mutation is 4 when peak iSAFE-score exceeds 0.1 versus a median rank of 10 along with a longer tail, when peak iSAFE-score is below 0.1 (Fig. S14). Empirically computed p-values (online methods) on iSAFE indicate good performance when p-value < 1e-4 (Fig. 3C).
Not surprisingly, iSAFE performance deteriorates when the favored mutation is fixed, or near fixation ($\nu > 0.9$ in Fig. 3A). To handle this special case, we include individuals from non-target populations. For a mutation, define the Maximum Difference in Derived Allele Frequency score (MDDAF) as the difference

$$\text{MDDAF} = D_T - \min(D_{NT}) \ ,$$  

where $D_T$ is the derived allele frequency in the target population and $\min(D_{NT})$ is the minimum derived allele frequency over all known non-target populations. Simulations of human population demography under neutral evolution (Fig. S15), shows $P(\text{MDDAF} > 0.78 | D_T > 0.9) = 0.001$ (see Fig. S16). Therefore, when we observe the rare event of high frequency mutations in target ($D_T > 0.9$) with MDDAF > 0.78, we add random outgroup samples to the data to constitute 10% of the data (online methods). In testing on the phase 3 of 1000 genome project (1000GP) data, we chose outgroup samples from non-target 1000GP populations. The addition of outgroup samples using the MDDAF criterion was tested in extensive simulations. While the performance did not change for $\nu < 0.9$, it dramatically improved for high frequencies, including when the favored mutation was fixed in the target population (Fig. 3A).

In testing instances of known human selective sweeps in 1000GP data, we note that performance is difficult to characterize due to many complicating factors. Multiple sweeps could be occurring in response to different selection events, including background selection in the same region, or polygenic selection may dilute the selection signal at any one locus. Moreover, the favored mutation is known unambiguously in only a few instances. We looked for genes/regions that showed the signature of a selective sweep in one of the 1000GP sub-populations, and had additional evidence pointing to the favored mutation. We identified 22 genes with some evidence, but only 8 ‘well characterized’ cases that presented irrefutable support for the favored mutation (Supp. Table S1).

We used iSAFE to rank all variants (~21,000) in a 5Mbp region surrounding the gene. Among the 8 well characterized cases, (Fig. 3B), iSAFE ranked the candidate mutations as 1 in 5 cases: SLC24A5, LCT, EDAR, ACKR1, TLR1; and, it assigned ranks 2 (ABCC1), 4 (HBB), and 13 (G6PD) in others. In almost all cases, we observed high iSAFE-scores ($\geq 0.1$). The spatial distribution of iSAFE-scores show a single, clear, peak in all 8 cases (Fig. 3F-M), in contrast to the iHS signal (Fig. 3D,E).

We checked to see if the other 14 regions under selection showed a strong iSAFE signal. In 3 of the 14 regions (FUT2, F12, ASPM), we only observed weak signals, and did not make a prediction (peak iSAFE < 0.027, p-val $\geq 0.008$), although we do see a strong iSAFE peak 1.3Mbp away from the ASPM gene (Fig. S24D). In other regions, iSAFE ranked the candidate mutations as 1 in the SLC45A2/MATP (CEU), MC1R (CHB+JPT), and ATXN2-SHB3 (GBR) genes, and 7, 8, and 12 in PSCA (YRI), ADH1B (CHB+JPT), and PCDH15 (CHB+JPT) genes, respectively. In each case, the iSAFE-scores were high with the exception of PSCA (peak iSAFE = 0.04, p-val = 2.4e-3, online methods).

The other 5 putative selected regions are interesting in that the top-ranked iSAFE mutations had high scores, but were distinct from the reported candidate mutations. Many of these genes
are involved in pigmentation, determining, skin, eye, and hair color. For example, the Tyrosinase (TYR) gene, encoding an enzyme involved in the first step of melanin production, is considered to be under positive selection with a nonsynonymous mutation rs1042602 as a candidate favored variant. A second intronic variant, rs10831496, in GRM5, 396kbp upstream of TYR, has been shown to have a strong association with skin color. In contrast, iSAFE ranks mutation rs672144 at the top. Interestingly, this variant was the top ranked mutation not only in CEU (iSAFE = 0.48, p-val ≤ 1.3e-8), but also in EUR, EAS, AMR, and SAS (iSAFE > 0.5, p-val ≤ 1.3e-8; Fig. S17). The result is consistent with the signal of selection being observed in all populations except AFR. It may not have been previously reported because it is near fixation in all populations of 1000GP except for AFR (Fig. S17H). We plotted the haplotypes carrying rs672144 and found that two distinct haplotypes carry the mutation, both remaining high frequency, maintained across a large stretch of the region, suggestive of a soft sweep with standing variation (Fig 4). A similar analysis applied to genes TRPV6, KITLG, OCA-HERC2 (Fig. 3R-T), where in each case, the top iSAFE mutations were identical across all non-African populations (online methods), and supported an out-of-Africa onset of selection. In the one remaining gene (CYP1A2/CSK; Fig. 3U), the top ranked iSAFE mutation rs2470893 was previously found significant in a genome wide association study, and was tightly linked to the candidate mutation. To summarize, iSAFE analysis ranked the candidate mutation among the top 13 in 14 of the 22 loci, did not show a strong signal in 3, and identified plausible alternatives in the remaining 5.

Discussion

The identification of the favored allele in a selective sweep is a long-standing computational problem in population genomics. Our results suggest that an understanding the coalescent structure of a region under a selective sweep can indeed pinpoint the favored mutation. iSAFE was designed to work in regimes where the selection strength is high, and there is a single favored mutation. However, its performance remains robust to a range of simulation parameters, including a wide range of initial frequencies (standing variation), and the frequency of the favored mutation at the time of sampling.

An important challenge was that regions undergoing a selective sweep also present a signal far away from the favored mutation, making it harder to pinpoint the favored mutation. The iSAFE technique, motivated by boosting with weak classifiers, exploits the soft shoulders. We observe when a true favored mutation is inserted into a shoulder region, it gets higher SAFE-scores on the average, in contrast to the insertion of a hitchhiking mutation. iSAFE uses this idea to rank mutations according to the weighted sum of their SAFE-scores in all windows.

We also use a cross-population technique in a limited manner by using the frequency differential of mutations in high frequency scenarios to get representative non-carrier haplotypes in the sample. Our future work will be aimed at seeing if the cross-population signal can be further improved, and if we can identify multiple favored loci within a region. Finally, we use only population based methods, and future work will seek to integrate these techniques with a functional analysis of
mutations.

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References


Figures

Figure 1: Illustration and Performance of the SAFE method. (A) The HAF score for haplotype \( h \) is the sum of the derived allele counts of the mutations on \( h \). (B) Carriers of the favored mutation have higher fraction of the total HAF score of the sample (high \( \phi \)) and lower number of distinct haplotypes compared to non-carriers (low \( \kappa \)). (C) Schematic of a no-recombination genealogy under a selective sweep. The mutations can be categorized as ‘non-carrier’ (gray), ‘ancestral to favored’ (green) arising prior to the favored mutation, and ‘descendant to favored’ (blue) that arise on haplotypes carrying the favored mutations but after the favored mutation, and the favored mutation itself (red). (D, E) Simulations showing \( \phi \) versus \( \kappa \) values for each variant in a neutral evolution and a selective sweep with default parameters. The joint-distribution of \( \phi \) and \( \kappa \), in a selective sweep, changes in a dramatic but predictable manner that separating out non-carrier (gray), descendant (blue), and ancestral (green) mutations from the favored (red) mutations. The SAFE score computes a normalized difference of the two statistics. (F) Performance (favored mutation rank) of SAFE vs. iHS for hard sweep \( (\nu_0 = 1/N = 1/20000) \) and soft sweep \( (\nu_0 > 1/N) \) on 50kbp window with 1000 simulations per bin.
Figure 2: iSAFE. (A) SAFE performance (rank distribution of favored mutation) as a function of window-size. The dashed line represents median rank, and decays for large windows while (B) iSAFE is robust to increase in window size. (C) The $\Psi(e, w)$ matrix, with $e \in S_1$, for a 5Mbp region around LCT gene in FIN population shows that the ‘shoulder’ of selection can extend for a few Mbp. The blue circle shows the location of the known favored mutation rs4988235. (D) iSAFE Illustration. The star, triangle, and circle denote the favored, ancestral, and descendant mutations, respectively. In its own window $w_1$, the triangle has a small number of distinct haplotypes. However, when inserted into $w_2$, the number of distinct haplotypes increases (increased $\kappa$). Analogously, when the circle is inserted in $w_2$, it reduces the proportion of total HAF-score (lower $\phi$). In contrast, the star (favored) mutation, retains high $\phi$ and low $\kappa$ when inserted in any windows. (E, F) Performance of iSAFE vs. iHS in a 5Mbp region simulation. (E) Cumulative Distribution Function (CDF) denotes the fraction of samples assigning a specific rank or lower. (F) The dashed (solid) lines provide the median (mean) rank as a function of favored allele frequency.
Figure 3: iSAFE Performance. (A) iSAFE performance upon addition of outgroup samples. No deterioration is seen for low frequencies of the favored variant, but iSAFE performance improves dramatically when favored mutation is near fixation or fixed. (B) iSAFE rank of candidate mutations (vs. iHS) for 8 well-characterized selective sweeps (Table S1). (C) Rank of favored mutation as a function of iSAFE-score (Bottom x-axis) or p-value (top x-axis). Each gray dot represents a sample drawn from simulations using a wide range of selection coefficients. The performance deteriorates for iSAFE-scores below 0.1. (F-M) iSAFE score plots on 8 well-characterized selective sweeps, with (D,E) iHS scores plotted for comparison for LCT and EDAR. Red stars represent the favored mutation in well-characterized selective sweep. The x-axis denotes location in cM. (N-U) iSAFE-scores on regions under selection. Top ranked iSAFE candidates are marked by blue squares when they match putative favored mutations, while green circles represent new favored mutations suggested by iSAFE. All data-sets were chosen by taking a 5Mbp window around the putative selected region, unless one side reached the telomere or centromere.
Figure 4: **The GRM5-TYR haplotype structure.** The mutation rs672144 is ranked first by iSAFE and very well separated from rest of the mutations in 5Mb around it, in all non-African populations with high confidence (iSAFE >0.5, \( p\)-val\( \ll \)1.3e-8). (A) Haplotype plot with core mutation rs672144 on all 2504 \( \times \) 2 haplotypes of 1000GP. This plot shows carrier haplotypes of mutation rs672144 are conserved over a longer span than haplotypes in non-carriers which is a signal of selection\(^8\). (B) Global frequencies of carrier haplotypes of mutation rs672144 (red, blue) and non-carrier haplotypes (gray). The evidence is consistent with an out of Africa selection on standing variation (soft sweep) with mutation rs672144 being the favored variant.
Online Methods

iSAFE: Input and Output. Consider a sample of phased haplotypes in a genomic region. We assume that all sites are biallelic and polymorphic in the sample. Thus, our input is in the form of a binary SNP matrix with each row corresponding to a haplotype and each column to a mutation, and entries corresponding to the allelic state, with 0 denoting the ancestral allele, and 1 denoting the derived allele. The output is a non-negative iSAFE-score for each mutation, according to its likelihood of being the favored variant of the selective sweep.

The Haplotype Allele Frequency (HAF-)score. The HAF score for haplotype \( h \) is the sum of the derived allele counts of the mutations on \( h \). Define the SNP matrix \( M \) such that, \( M_{h,e} = 1 \) if haplotype \( h \) carries the derived allele of SNP \( e \), and 0 otherwise. The Haplotype Allele Frequency (HAF) score of haplotype \( h \) defined in Eq. (1) as:

\[
\text{HAF}(h) = \sum_e M_{h,e} \sum_{h'} M_{h',e} = \sum_{h'} [M \cdot M^T]_{h,h'},
\]

where \( \sum_{h'} M_{h',e} \) is derived allele count for SNP \( e \), and \( [M \cdot M^T]_{h,h'} \) is number of shared derived alleles (mutations) between haplotypes \( h \) and \( h' \) (see Fig. 1A). The HAF score is shown to be very helpful in predicting carriers of ongoing selective sweeps without knowledge of the favored allele[6].

SAFE: Selection of Allele Favoured by Evolution. For each SNP \( e \), define \( \phi \) as:

\[
\phi(e) = \frac{\sum_h [M_{h,e} \cdot \text{HAF}(h)]}{\sum_h \text{HAF}(h)},
\]

that is sum of HAF scores of carriers of the derived allele \( e \) (\( \sum_h [M_{h,e} \cdot \text{HAF}(h)] \)), divided by sum of HAF scores of all haplotypes in the sample (\( \sum_h \text{HAF}(h) \)). For each SNP \( e \), we define \( \kappa \) score as:

\[
\kappa(e) = \frac{\left| \bigcup_h \{M_{h,e} \cdot \text{HAF}(h)\} \right|}{\left| \bigcup_h \{\text{HAF}(h)\} \right|} - 1,
\]

that is the number of distinct non-zero values in HAF scores of SNP \( e \) carriers, divided by number of distinct values in HAF scores of all haplotypes in the sample population. For each SNP \( e \), we define SAFE score as:

\[
\text{SAFE}(e) = \frac{\phi(e) - \kappa(e)}{\sqrt{f_e(1-f_e)}},
\]

where \( f_e \) is the derived allele frequency of SNP \( e \).

Empirical analysis on simulation data shows that for a neutrally evolving population, \( \phi \) and \( \kappa \) are biased estimators of derived allele frequency \( f \) (Fig. S1) and \( \lambda f(1-f) \) is a biased estimator for variance of \( (\phi - \kappa) \), where \( \lambda \) is a positive constant. Consequently, we assume that the distribution of
the SAFE score of derived alleles in a neutrally evolving population is approximated by a *Gaussian* distribution with mean 0 and unknown variance $\lambda$ (see Fig. S2).

For a population undergoing a positive natural selection, $\phi$ over estimate, and $\kappa$ under estimate (Fig. 1F) the favored allele frequency ($\nu$). Therefore, we expect the distribution of ($\phi - \kappa$) for the favored allele to be skewed in positive direction.

Performance of SAFE score for detecting the favored variant on a small window is promising (See Figs S3, S4, S7, S5, S6); but the performance decays in larger windows (Fig. S7); because in larger windows most of the haplotypes become unique and $\kappa$ estimate $f$ correctly, even for favored mutations of selective sweeps, while we expect it to underestimate the $f$ for the favored mutations. Consequently, the estimator $\kappa$ is no-longer useful for pinpointing the favored mutation.

**iSAFE: integrated SAFE for large regions.** We devise iSAFE-score by extending the SAFE score to boost the performance in larger windows. We apply the SAFE score, as a kernel, on overlapping sliding windows. Define $S$ as the set of all SNPs, $W$ as the set of all sliding windows. Define the score $\alpha$ of window $w \in W$ as:

$$\alpha(w) = \frac{\sum_{e \in S_1} \Psi_{e,w}}{\sum_{w \in W} \sum_{e \in S_1} \Psi_{e,w}}, \quad (S5)$$

where $\Psi_{e,w}$ is the SAFE score of SNP $e \in S$ assuming it is in window $w \in W$ if it is positive, 0 otherwise; and $S_1$ is the union of first rank mutations of all $w \in W$. Define the score iSAFE of SNP $e \in S$ as:

$$iSAFE(e) = \sum_{w \in W} \Psi_{e,w} \cdot \alpha(w). \quad (S6)$$

**Maximum Difference in Derived Allele Frequency (MDDAF).** We have shown that iSAFE is successful in pinpointing the favored variant in an ongoing selective sweep. When the favored mutation is near fixation ($\nu > 0.9$), iSAFE performance decays and when the favored variant is fixed ($\nu = 1$), iSAFE cannot detect the favored mutation because it is no longer a variant (Fig. 3A). For the purpose of pinpointing the favored mutation in a fixed selective sweeps we add random sample from non-target population (outgroup) to the target population to constitute 10% of the sample.

To minimize the noise added to the data with random outgroup samples, we devise a simple method to decide whether to use outgroups or not. Our score is motivated by the work of Grossman et al. (2010) who introduced the $\Delta$DAF score of a mutation as $\Delta$DAF = $D_T - D_{NT}$, where $D_T$ is the derived allele frequency in the target population and $D_{NT}$ is the *average* derived allele frequency in non-target populations. As it is possible that some of the non-target populations are also under selection, choosing the average derived allele frequency may lower $\Delta$DAF, and weaken the signal of selection. Instead we define the Maximum Difference in Derived Allele Frequency (MDDAF) score as:

$$MDDAF = D_T - \min(D_{NT}), \quad (S7)$$
where, $D_T$ is the derived allele frequency in the target population and $\min(D_{NT})$ is the minimum derived allele frequency over all non-target populations.

**Adding Outgroup Samples.** Simulation of human population demography under neutral evolution (Fig. S15), shows $P(MDDAF > 0.78|D_T > 0.9) = 0.001$ (Fig. S16) making it a rare event to have high MDDAF score even when the frequency is high in the Target population. Therefore, when there is a high frequency mutation ($D_T > 0.9$) with MDDAF $> 0.78$ in the target population, we add random outgroup samples to the data to constitute 10% of the data. For analysis on real data, where we looked at 1000GP populations, we randomly selected outgroup samples from non-target populations of 1000GP.

In Fig. 3A, we compared the performance of iSAFE with or without having the option of using outgroup samples; we simulated 5Mbp of human genome based on the human demography model described in Fig. S15. The selection happens in a random time after the out of Africa in EUR population (as the target population). When the onset of selection is before split of EUR and EAS, both (EUR and EAS) are under selection. When we have random sample option, we use the MDDAF criterion to decide whether we should use random sample or not. In case of adding random sample, we add a random subset of individuals from EAS+AFR to constitute 10% of the data (200 haplotypes from EUR and 22 from EAS+AFR).

The performance of iSAFE for sweeps with $\nu < 0.9$ did not change with or without having outgroup sample option (Fig. 3A). When frequency of the favored mutation is near fixation ($\nu > 0.9$) having the outgroup sample option is helpful and increase the performance of the iSAFE. When the sweep is fixed ($\nu = 1$), iSAFE is no longer capable of detecting the favored mutation without having outgroup samples because the favored mutation is no longer a variant in the target population. However, with the outgroup sample option, iSAFE can successfully pinpoint the Favored mutation even in a fixed selective sweep (see Fig. 3A).

**Simulations.**

Neutral and sweep samples were generated using the simulator msms\(^7\). By default, simulated populations are haploid with sample size of $n = 200$ haplotypes from a larger effective population of $N = 20000$ haplotypes, each of length $L$, with default value 50kbp for SAFE and 5Mbp for iSAFE. For human populations, a mutation rate of approximately $\mu = 2.5 \cdot 10^{-8}$ mutations per bp per generation\(^12\), and a recombination rate of approximately $r = 1.25 \cdot 10^{-8}$ per bp per generation\(^14\) have been proposed. For SAFE simulations, we used a scaled mutation rate $\theta = 2\mu N = 1$ mutations per kbp per generation and scaled recombination rate $\rho = 2rN = 0.5$ crossovers per kbp per meosis to approximate human rates. The rates were scaled linearly by $L$. In the case of positive selection the default scaled selection strength of the favored allele was set to $Ns = 500$, with the favored mutation located at a random position uniformly distributed on the range $[1, L]$. The default value for favored mutation starting frequency $\nu_0 = 1/N$ (hard sweep), and the frequency of the favored mutation ($\nu$) at the time of sampling is a random value uniformly distributed on the range $[0.1, 0.9]$. We simulated demography of AFR, EUR, EAS populations with parameter shown in the Fig. S15.
based on a popular demographic model of human population. We used the default parameters for all simulations unless otherwise stated.

**Empirical \(p\)-val computation.**

We applied iSAFE on a neutrally evolving simulated population with window size 5Mbp, based on European demography shown in Fig. S15. A \(p\)-value was calculated based on empirical distribution of iSAFE on these simulated populations. We limited the number of samples to \(\sim 74,800,000\) for efficiency, and this allows us to get a \(p\)-value as low as 1.34e-8 for iSAFE-score 0.304. Scores higher than this cut-off are considered to have \(p\)-value \(< 1.34e-8\).

**Results on selective sweeps in human populations**

**Well characterized selective sweeps.** We examined 8 well characterized selective sweeps with strong candidate mutation. These genes are LCT, SLC24A5, TLR1, EDAR, ACKR1/DARC, ABCC11, HBB, and G6PD. iSAFE results for these genes are summarized in Fig. 3 and Table S1.

We also examined 14 other regions reported to be under selection with one or more candidate favored mutations.

**Pigmentation genes.**

**SLC45A2/MATP.** This region is involved in human pigmentation pathways and is a target of selective sweep in European population. A nonsynonymous mutation rs16891982 is associated with light skin pigmentation and is believed to be the favored variant. This mutation is also ranked first by iSAFE out of \(\sim 21,000\) mutations (5Mbp) in CEU population with a significant score (see Fig. 3N, iSAFE=0.32, \(p\)-val\(< 1.3e-8\)). This mutation is almost fixed in European; frequency in AFR, EAS, SAS, AMR, and EUR is 0.04, 0.01, 0.06, 0.45, and 0.94, respectively.

**MC1R.** The MC1R gene is implicated in many skin color phenotypes, including red hair, fair skin, freckles, poor tanning response and higher risk of skin cancer. It is a target of positive selection in East Asian populations, with a non-synonymous mutation (rs885479) suggested as a candidate favored mutation. This mutation is ranked first by iSAFE in CHB+JPT (see Fig. 3P, iSAFE =0.24, \(p\)-val \(= 1.4e-6\)) out of \(\sim 16,000\) mutations (2.8Mbp). The putative selected region is 300kbp away from the telomere of chromosome 16.

**GRM5-TYR.** The Tyrosinase (TYR) gene, encoding an enzyme involved in the first step of melanin production is present in a large region under selection. A nonsynonymous mutation rs1042602 in TYR gene is reported as a candidate favored variant. A second intronic variant rs10831496 in GRM5 gene, 396kbp upstream of TYR, has been shown to have a strong association with skin color.
In contrast, iSAFE ranks mutation rs672144 as the top candidate for the favored variant region out of \( \sim 22,000 \) mutations (5Mbp). This variant was the top ranked mutation not only in CEU (iSAFE = 0.48, \( p \)-val\( \ll 1.3e^{-8} \)), but also the top ranked mutation for EUR, EAS, AMR, and SAS (see Fig. 3Q and Fig. S17). The signal of selection is strong in all populations (iSAFE >0.5, \( p \)-val\( \ll 1.3e^{-8} \) for all) except AFR, which does not show a signal of selection in this region. It may not have been reported earlier because it is near fixation in all populations of 1000GP except for AFR (\( f = 0.27 \)), as seen in Fig. S17G. We plotted the haplotypes carrying rs672144 and found (Fig. 4) that two distinct haplotypes carry the mutation, both with high frequencies maintained across a large stretch of the region, suggestive of a soft sweep with standing variation.

The previously suggested candidates rs1042602, rs10831496 are fully linked to rs672144 (Fig. S18), but not to each other. The EUR haplotypes can be partitioned into 4 clusters (Fig. S18). Each of the 4 haplotypes show high homozygosity, suggestive of selection. However, rs1042602 can only explain the sweep in clusters C1+C2. rs10831496 can only explain C1+C3. Only rs672144 explains all 4 clusters, providing a simpler explanation of selection in this region. GTEx eQTL analysis on TYR gene for the tissue ‘Skin - Sun Exposed (Lower leg)’ showed \( p \)-value 0.61 for rs1042602, \( p \)-value 0.15 for rs10831496, and \( p \)-value = 0.08 for rs672144. While the \( p \)-value does not rise to a level of significance due to sample size issues, it is indicative of a regulatory function for the mutation.

**OCA2-HERC2.** This region is suggested as a target of selection in European, and several mutations in this region are associated with hair, eye, and skin pigmentation. For example, rs12913832 is considered to be the main determinant of iris pigmentation (brown/blue) and is also associated with skin and hair pigmentation and the propensity to tan. rs1667394 is also linked to blond hair and blue eyes. Some other mutations, many fully linked, (rs4778138, rs4778241, rs7495174, rs1129038, rs916977) are also associated with blue eyes. This region is also suggested to be a target of selection in East Asia with rs1800414 suggested as a candidate for light skin pigmentation in that population. We applied iSAFE on this region to all 1000GP super-populations.

iSAFE selected a single variant rs1448484 in OCA2 (with high confidence, \( p \)-val\( <1.34e^{-8} \) for EUR, EAS, AMR and \( p \)-val\( =2.13e^{-6} \) for SAS) as the favored variant in all 1000GP populations (EUR, EAS, SAS, AMR) except for AFR that showed no signal of selection in this region (see Fig. S19 and Fig. 3P). This variant is close to fixation in all populations except for AFR, where \( \nu = 20\% \) (see Fig. S19F). iSAFE result along with the frequency pattern of the top ranked variant, suggests an out of Africa selection, probably on light skin color, on this region. The other candidate variants are all ranked high, and tightly linked with the top-ranked variant (Table S2).

**KITLG.** This genomic region has been linked to skin pigmentation in European and East Asian populations, and shows a strong signature of selective sweep on regulatory regions surrounding
the gene in all non-African populations[21], with a candidate variant rs642742, that is associated
with skin pigmentation[25].

iSAFE analysis identified the same mutations gaining the top rank in multiple populations
(Fig. S20). Top rank mutations in EUR, SAS, EAS, and AMR populations are shown in
Table S3. The top ranked mutation in EUR and CEU populations (rs405647) was ranked
1, 2, 3 in AMR, SAS, and EAS, respectively, and is tightly linked to rs642742 ($D' = 0.92$).
Mutation rs661114 is ranked 2 in EUR, 5 in CEU, 6 in SAS, and 20 in AMR, and lies in a
region with H3K27 acetylation that is associated with enhanced expression.

**TRPV6.** This region has been reported a target of selection in CEU population[22]. TRPV6 is
involved in calcium absorption. It has been suggested that “Individuals with lighter skin pig-
mentation might have produced too much 1,25-dihydroxyvitamin D, resulting in an increased
intestinal Ca2+ absorption. Thus, to reduce the risk of absorptive hypercalciuria with kidney
stones, the derived haplotype would have spread only among individuals with lighter skin pig-
mentation”[23]. iSAFE suggests 10 strongly linked mutations located along a 9kbp region
located 84kbp downstream of TRPV6 (see Fig. S22). These mutations are ranked in the top
10 in all non-African populations (Table S5). There is no signal of selection in this region in
AFR. The pattern of selection in this region in global population along with the confidence
and consistency of iSAFE results in all non-African populations is consistent with an out of
Africa selection on this region with the favored mutation being near fixation in all non-African
populations (Fig. S21).

**Population specific selection: East Asian.**

**PCDH15.** This gene plays a role in development of inner-ear hair cells and maintaining retinal
photoreceptors and is reported to be under selection in East Asian and a nonsynonymous
mutation rs4935502 is proposed to be the favored variant[4]. This mutation is ranked 12 by
iSAFE in CHB+JPT (see Fig. S24A, iSAFE =0.45, p-val<1.34e-8). All top mutations are
highly linked.

**ADH1B.** "The ADH1B gene encodes one of three subunits of the Alcohol dehydrogenase (ADH1)
protein, a major enzyme in the alcohol degradation pathway that catalyzes the oxidization
of alcohols into aldehydes.” This region is a target of positive selection in East Asian popu-
lation[27]. A non-synonymous mutation in this gene is associated with Alcohol dependence[27].
We tested this gene in CHB+JPT populations. iSAFE rank, in 2Mbp around ADH1B gene,
for the candidate mutation (rs1229984) is 8 (see Fig. S24B). The top rank mutation is an up-
stream mutation (rs3811801) 5kbp upstream of the candidate mutation rs1229984 and highly
linked to it ($D' = 0.99$). The second rank mutation (rs284787) is a 3′-UTR of ADH7 which is
shown to be associated with Upper Aerodigestive Tract Cancers in a Japanese Population[28].
Population specific selection: UK

The UK Biobank project was recently investigated for regions under selection. The regions were reported as a target of a recent selection by analyzing the structure of UK Biobank and Ancient Eurasians. We applied iSAFE on GBR (British in England and Scotland) population in 1000GP to check if the favored mutation could be confirmed.

**ATXN2-SH2B3.** Galinsky et al. proposed a nonsynonymous mutation (rs3184504) as a candidate that is associated to blood pressure. We tested this region in GBR population of 1000GP. This candidate mutation is jointly ranked first with two other mutations rs7137828, rs7310615 (see Fig. 3O, iSAFE = 0.27, p-val=1.6e-7). rs7137828 is an intronic mutation in ATXN2 that is associated with Primary Open Angle Glaucoma that is a leading cause of blindness worldwide. The other first rank mutation (rs7310615) is associated with blood expression levels of SH2B3. Surprisingly, all of the top 10 mutations, ranked by iSAFE have a known association to a phenotype (Table S4), and are highly linked (Fig. S23).

**CYP1A2/CSK.** We tested a 5Mbp region around these genes in GBR population of 1000GP. The proposed mutation rs1378942 by with frequency 0.69 in GBR population is ranked 89 by iSAFE (iSAFE = 0.13, p-val=7.0e-5). The top-ranked mutation rs2470893 (Fig. 3U, iSAFE = 0.16, p-val=2.7e-5) is between CYP1A1 and CYP1A2 with frequency 0.40 in GBR and is associated with Caffeine metabolism. rs2470893 and rs1378942 are in a strong LD ($D' = 0.91$).

**FUT2.** The signal of selection on 5Mbp around this region in GBR population is very weak (Fig. S24E), with peak iSAFE = 0.026, p-val=0.009. There is a very weak peak in 400kbp around FUT2 gene (chr:49077276-49475876). The stop gained mutation rs601338 proposed as a candidate mutation by is ranked 4 (p-val=0.1).

**F12.** The signal of selection on 5Mbp around this region in GBR population is very weak (Fig. S24F, peak iSAFE = 0.027, p-val=0.008). The proposed mutation rs2545801 has a very weak signal (p-val=0.2).

**Other genes**

**PSCA.** This gene has been reported as a target of selection in YRI population. A 5'UTR mutation rs2294008 proposed as a candidate favored mutation in this region that is associated with urinary bladder and gastric cancers. The signal of iSAFE in 5Mbp around this gene in YRI population is weak (see Fig. S24C, peak iSAFE = 0.04, p-val=2.4e-3). The proposed mutation rs2294008 is ranked 7 in 5Mbp region surrounding this region. The local rank in 400kbp around this gene is joint-first with 8 other mutations including rs2976392 which is also associated with diffuse-type gastric cancer. Other mutations are rs2978979, rs2920279, rs2978980, rs2920282, rs2294010, rs2717562, rs2978982. This 9 mutation are fully linked in
YRI population in a 20kbp region that cover PSCA from upstream regulatory region to its downstream (chr8:143757286-143776668, GRCh37/hg19).

**ASPM.** This gene is reported to be a target of weak selection in GBR population\(^{22}\). The signal in 2Mbp around this gene is very weak (see Fig. S24D, peak-iSAFE = 0.025, \(p\)-val=0.01). The proposed mutation rs41310927 has a very weak signal (\(p\)-val=0.4). However, we do see a strong iSAFE signal 1.3Mbp away from the ASPM gene.
Figure S1: $\kappa$ and $\phi$ as estimators of $f$. Empirical analysis, with 10,000 neutrally evolving population (about 3 million SNPs) with default parameter set, shows that $\phi$ and $\kappa$ are (biased) estimators of allele frequency $f$.

Figure S2: **Empirical SAFE distribution.** (A) SAFE score Probability Density Function (PDF) of 10,000 neutrally evolving population (about 3 million SNPs) with default parameter set. (B) Quantiles of the SAFE score against the quantiles of the Normal distribution for the same data in part A. The coefficient of determination ($R^2 = 0.9997$) for the QQ-plot shows that Gaussian distribution is a good approximation to the SAFE score distribution.
Figure S3: **Performance of SAFE score on hard and soft sweeps.** (A) Rank of the favored mutation for hard sweep ($\nu_0 = 1/N = 1/20000$) and soft sweep ($\nu_0 > 1/N$) in 1000 simulations per bin on 50kbp window with selection strength ($N_s = 500$) and default values for other simulation parameters. The line with large dashes represents the median rank. (B) Rank of the favored mutation as a function of Origin Count (number of ancestors of carriers of favored Allele at the onset of selection pressure) for the same data as in A. Origin Count of hard sweep is always 1.
Figure S4: **Performance of SAFE score with different selection strength.** (A, B) Rank and rank percentile of the favored mutation as a function of selection strength \( (Ns) \) in 1000 simulations per bin on 50kbp window with default values for other simulation parameters. The line with large dashes represents the median rank.
Figure S5: **Performance of SAFE score with different sample size.** (A, B) Rank and rank percentile of the favored mutation as a function of sample size in 1000 simulations per bin with selection strength \((N_s = 500)\) and default values for other simulation parameters. The line with large dashes represents the median rank.

Figure S6: **Performance of SAFE score in a model of European demography.** White represents the result for a fixed size population model with default parameters and gray represents a model of human demography for EUR population. The model and all the parameters used are described in Fig. S15. The onset times of selection was post-bottleneck (23 kya-current) epochs. 1000 samples per bin were simulated with default values for simulation parameters not assigned in Fig. S15.
Figure S7: **Performance of SAFE score with different window size.** (A, B) Rank and rank percentile of the favored mutation as a function of window size in 1000 simulations per bin with selection strength ($N_s = 500$) and default values for other simulation parameters. The line with large dashes represents the median rank.
Figure S8: **iSAFE and Window Size.** Performance of iSAFE measured by rank of the favored variant and the distance of the favored variant from the peak in 1000 simulations per bin. The line with large dashes represents the median rank.

Figure S9: **iSAFE and Position of the Favored Mutation.** iSAFE rank of the favored mutation on 5Mbp regions with different position of the favored mutation. Each bin includes 1000 simulations with the position of the favored mutation selected from [0.5Mbp, 1.5Mbp, 2.5Mbp]. The dashed (dotted) line represents median (upper quartile). This result shows iSAFE is robust to the position of the favored mutation.
Figure S10: **iSAFE vs. iHS in hard sweep.** Performance of iSAFE vs. iHS, in hard sweep ($\nu_0 = 1/N = 1/20000$), measured by rank of the favored variant and the distance of the favored variant from the peak in 5000 simulations on 5Mbp region. **(A,B)** Solid (dashed) lines represent the mean (respectively, median) value of the favored allele rank (Panel A), and of the distance of the iSAFE peak from the favored allele (Panel B). **(C)** For any rank $r$ on the X-axis, the $y$-intercept represents the proportion of samples where the favored allele had rank $\leq r$. **(D)** For any distance $d$ on the X-axis, the $y$-intercept represents the proportion of samples where the favored allele had distance $\leq d$ from the iSAFE peak.
Figure S11: **iSAFE vs. iHS in soft sweep.** Performance of iSAFE vs. iHS, in in hard sweep ($\nu_0 = 1/N = 1/20000$) and soft sweep ($\nu_0 > 1/N$), measured by rank of the favored variant and the distance of the favored variant from the peak in 5000 simulations on 5Mbp region.
Figure S12: **iSAFE Demo I.** iSAFE on 5Mbp region with different selection strength, \( Ns \in [0, 100, 200, 500, 1000] \). Left panels show the \( \Psi^1 \) matrix. Middle panel shows the iSAFE-score as a function of the variant position. Right panel show the derived allele frequency as a function of iSAFE.
Figure S13: **iSAFE Demo II.** iSAFE on 5Mbp region with position of the favored mutation selected from range [2.5Mbp, 5Mbp] and with different favored allele frequency $\nu$. 
Figure S14: **Peak iSAFE**. Empirical analysis, with 5000 simulations on 5Mbp region with a wide range of selection strength \((N_s \in [10, 50, 100, 200, 300, 400, 500, 1000])\), shows difference in performance of iSAFE beyond a score threshold of 0.1 for peak value of iSAFE (see Fig. 3C).
Figure S15: A model of human demography described by Gravel et al. (2011)\textsuperscript{15} Fig. 4, Table 2. The model assumes an out-of-Africa split at time \( T_B \), with a bottleneck that reduced the effective population from \( N_{Af} \) to \( N_B \), allowing for migrations at rate \( m_{Af-B} \). The African population stays constant at \( N_{Af} \) up to the present generation. The model assumes a second split between European and Asian populations at time \( T_{EuAs} \), with a bottleneck reducing the Asian and European populations to \( N_{As0} \) and \( N_{Eu0} \) respectively. The bottleneck was followed by exponential growth at rates \( r_{As} \) and \( r_{Eu} \), as well as migrations among all three sub-populations, leading to current populations from which East Asian (EAS), European (EUR), and Africans (AFR) individuals were sampled.
Figure S16: **Maximum Difference in Derived Allele Frequency (MDDAF)**. We simulated 25,000 instances of AFR, EUR, and EAS populations, based on a demographic model described in Fig. S15. We used default values for simulation parameters not assigned in the Fig. S15. 

**A)** The MDDAF score of mutations as a function of derived allele frequency in the target population $D_T$. 

**B)** Distribution of the MDDAF score for mutations with $D_T > 0.9$. 

**C)** P-value of the MDDAF score for mutations with $D_T > 0.9$. The dashed-red lines represent the value $0.78$, where MDDAF, given $D_T > 0.9$, has a p-value less than $0.1\%$. 

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Figure S17: **iSAFE on GRM5-TYR.** The mutation rs672144 is the iSAFE top rank mutation in all the population of 1000GP except African that doesn’t show any signal of selection in this region.
Figure S18: **GRM5-TYR SNP matrix.** Each row is a haplotype and each column is a variant in EUR populations of 1000GP. In total we have 1006 haplotypes. Carriers haplotypes of derived alleles of rs10831496, rs672144, and rs1042602, are shaded by red, green, and blue, respectively. For making the plot sensible, We removed low frequency SNPs $f_{EUR} < 0.2$ and SNPs that are near fixation in the whole 1000GP, $f_{1000GP} > 0.95$. 
Figure S19: **OCA2-HERC2.** The mutation rs1448484 is the iSAFE top rank mutation in all the population of 1000GP except African that doesn’t show any signal of selection in this region. rs12913832 is a candidate favored mutation for the selection in European, proposed by \(^9\). Table S2 provides iSAFE rank of some other candidate mutations associated with pigmentation.\(^{24}\)
Figure S20: KITLG. iSAFE top rank mutations (circles) and candidate mutation rs642742 (blue triangle) proposed by 25. See the Table S3 for more details.
Figure S21: **TRPV6.** 10 mutations (rs11772526, rs4725602, rs11763225, rs7796010, rs11762011, rs13239916, rs4145394, rs10808023, rs10808021, and rs4726591) are highly linked and are top 10 iSAFE candidate mutations in all the 1000GP populations except for AFR where there is no signals of selection.
Figure S22: TRPV6. Haplotypes of top 10 iSAFE mutations, and the proposed mutation (rs4987682) by [22], in 5Mbp around TRPV6 in 2504 × 2 haplotypes of 1000GP are shown. These mutations are sorted by their iSAFE rank from left to right. iSAFE top 10 mutations span a 9kbp region(chr7:142476441-142485399, GRCh37/hg19). White is derived and black is ancestral allele.

Figure S23: ATXN2-SH2B3. Haplotypes of top 20 iSAFE mutations in 5Mbp around ATXN2-SH2B3 in GBR population are shown. These mutations are sorted by their iSAFE rank from left to right. They span a 1.07Mbp region around ATXN2-SH2B3 region (chr12:111833788-112906415, GRCh37/hg19). White is derived and black is ancestral allele. Most of these mutations are associated to a phenotype (see Table S4).
Figure S24: **iSAFE on Targets of Selection.** iSAFE-scores on regions under selection. Putative favored mutation is shown in blue square when it is among iSAFE top rank mutations, and in blue triangle when the signal of selection is very weak.
## Table S1: iSAFE on 8 well characterized selective sweeps.

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Table S2: **OCA2-HERC2.** iSAFE rank of candidate mutations proposed by\[^{21}\] in 1Mb region around OCA2-HERC2 that are associated with eye, hair, and skin pigmentation.

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Table S3: **KITLG**. iSAFE rank of top mutations in 2 Mbp around KITLG gene. sorted by their Mean Reciprocal Ranks, calculated over EUR, SAS, EAS, AMR. Only those with Mean Reciprocal Rank greater than 0.1 are shown (the candidate mutation rs642742 proposed by\textsuperscript{25} is also reported in the last row). Frequency and iSAFE score for this region in all the 1000GP populations are provided in S20.

<table>
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<th>iSAFE Rank SAS</th>
<th>iSAFE Rank EAS</th>
<th>iSAFE Rank AMR</th>
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Table S4: **ATXN2-SH2B3.** iSAFE rank of top 20 mutations in GBR population of 1000GP in 5Mbp around ATXN2-SH2B3 region and their association to diseases.

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<th>Rank</th>
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<th>Function</th>
<th>GBR Frequency</th>
<th>Association</th>
<th>Reference</th>
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<td>missense</td>
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<td>Blood pressure and hypertension, Coronary artery disease, &amp; more</td>
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<td>0.5</td>
<td>Primary open-angle glaucoma</td>
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<tr>
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<td>Systemic lupus erythematosus, Retinal vascular caliber</td>
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<tr>
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<td>Body mass index</td>
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<tr>
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Table S5: **TRPV6.** iSAFE rank of top mutations in 5Mbp around TRPV6 gene. sorted by their Mean Reciprocal Ranks, calculated over EUR, SAS, EAS, AMR.

<table>
<thead>
<tr>
<th>ID</th>
<th>iSAFE Rank EUR</th>
<th>iSAFE Rank SAS</th>
<th>iSAFE Rank EAS</th>
<th>iSAFE Rank AMR</th>
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