Visualization of Pairwise and Multilocus Linkage Disequilibrium Structure Using Latent Forests

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Abstract

Linkage disequilibrium study represents a major issue in statistical genetics as it plays a fundamental role in gene mapping and helps us to learn more about human history. The linkage disequilibrium complex structure makes its exploratory data analysis essential yet challenging. Visualization methods, such as the triangular heat map implemented in Haploview, provide simple and useful tools to help understand complex genetic patterns, but remain insufficient to fully describe them. Probabilistic graphical models have been widely recognized as a powerful formalism allowing a concise and accurate modeling of dependences between variables. In this paper, we propose a method for short-range, long-range and chromosome-wide linkage disequilibrium visualization using forests of hierarchical latent class models. Thanks to its hierarchical nature, our method is shown to provide a compact view of both pairwise and multilocus linkage disequilibrium spatial structures for the geneticist. Besides, a multilocus linkage disequilibrium measure has been designed to evaluate linkage disequilibrium in hierarchy clusters. To learn the proposed model, a new scalable algorithm is presented. It constrains the dependence scope, relying on physical positions, and is able to deal with more than one hundred thousand single nucleotide polymorphisms. The proposed algorithm is fast and does not require phase genotypic data.

Introduction

Linkage disequilibrium (LD) refers to non-random associations of alleles at two or more loci, over the human genome [1,2]. LD is usually present at short-range, i.e., for distances less than 10 kb [3]. Nevertheless, long-range LD (i.e., LD with distances greater than 100 kb) [3], and LD between different chromosomes [4], are also observed. Analyzing the extent and distribution of LD represents a major topic in statistical genetics. For instance, LD plays a fundamental role in gene mapping: the observation of a large number of genetic markers over a chromosomal region ensures a precise localization of (non-observed) causal mutations. Based on this property, genome-wide association studies (GWASs) [5,6] aim to systematically localize causal loci over the genome using hundreds of thousands of single nucleotide polymorphisms (SNPs), an abundant and useful class of genetic markers. Beside gene mapping, LD pattern analysis offers deep insights into the understanding of human population history. Bottlenecks, natural selection and migrations are examples of evolutionary events which can be inferred using coalescent models [7].

At the interface between computer science and artificial intelligence, data mining (DM) is the process of extracting patterns from data [8]. DM helps formulate hypotheses worth testing and is complementary to more conventional statistics. Data visualization, a branch of DM, aims at providing efficient and intuitive tools to represent and summarize relevant information underlying data [9]. Data visualization has been successfully applied to bioinformatics [10]. The international HapMap project [3], and more recently the international 1000 Genomes project [11], have made considerable efforts to deeply characterize the genome sequence variation in human populations. In this context, the application of visualization methods in the analysis of LD patterns has been shown to be essential, most notably to reveal the complex so-called LD block structure [12]. The simplest but also the most popular method is the triangular heat map (THM) as implemented in Haploview [13]. The THM is the triangular matrix of pairwise dependences between genetic markers, in which the color shading indicates the LD strength in each matrix cell. The THM generally displays the Lewontin D′ or the squared correlation coefficient r2. Another dependence measure, the ratio of the D′ to the logarithm of odds (noted LOD), is used as a standard by Haploview. In the THM, LD blocks are visually apparent. Nevertheless, the THM has the drawback to only display pairwise dependences, thus providing a restricted view of multilocus patterns. Another popular approach consists in plotting the fine-scale map of recombination rates computed along the chromosomal sequence. For this purpose, PHASE [14], a coalescent-based method, can be used to estimate recombination rates between adjacent SNPs in the sequence. This approach helps find recombination hotspots and provides insight of the underlying block structure of LD, but leads to computational burden. More advanced techniques, such as those providing isometric blocks and bifurcation plots [15], or textile plots [16], can deal with multilocus LD. For instance, the algorithm used to draw a textile plot is closely related to principal component analysis. The textile plot strategy consists in assigning the optimal...
geometrical configuration to variables and data points in a low-dimensional linear space.

At the interface of graph and probability theories, probabilistic graphical models (PGMs) represent a powerful formalism to uncover complex networks of interactions. Thanks to their ability to capture (conditional) independences and dependences between variables, PGMs offer an accurate modeling of relationships between variables in an uncertain framework [17]. A PGM is a probabilistic model that relies on a graph representing conditional independences within a set of random variables. Essentially, this model provides a compact and natural representation of the joint probability distribution of the variable set. PGMs have been successfully applied to LD modeling, in particular for haplotype inference and association genetics [18–21]. Recently, Mourad et al. introduced forests of hierarchical latent class models (FHLCMs) to model genome-wide LD, together with a scalable algorithm, named CFHLC (Construction of Forests of Hierarchical Latent Class models), able to cope with $10^5$ variables and 2000 individuals [22,23]. FHLCMs will be described in details in the next section.

In this paper, we describe another attractive property of FHLCMs (beside LD modeling) as LD visualization tools. We advocate their use for: (i) short-range, (ii) long-range and (iii) chromosome-wide LD visualization. Most notably, these models provide a compact and interpretable view of LD for the geneticist, thanks to their hierarchical graphical nature and their latent variables (LVs). Moreover, the proposed method allows to visualize both pairwise and multilocus LD on a single display, and to tackle the fuzziness of LD block boundaries. Low-level LVs represent short-range LD and are interpreted as haplotype shared ancestry, whereas high-level LVs correspond to long-range LD and are seen as population structure or natural selection effects. We also define a new multilocus LD measure for the evaluation of LD strength inside FHLCM clusters. Tested on real datasets, our method has been shown to be a valuable tool for the geneticist, regarding both information summary and understanding of LD spatial structure.

A first version of CFHLC, the learning algorithm, requires the splitting of the genome in large fixed-size windows (100–600 SNPs) in order to process genome-scale data. This represents a severe drawback as the possible dependences between adjacent windows are missing, which blurs the analysis of long-range LD. Thus, for an adequate use of FHLCMs in LD visualization, we have developed a new version of the CFHLC algorithm (named CFHLC+). In this second version, the genome splitting is not requested anymore to resolve the scalability issue. LD modeling is now constrained by physical position of SNPs along the chromosome. This constraint, less drastic than genome splitting, actually corresponds to a sliding window approach. Fixing the sliding window size sufficiently large ($0.1–5 \text{ Mb}$) represents a reasonable strategy to capture long-range LD in the GWAS context.

This paper is organized as follows. Section Results illustrates our LD visualization approach within three experimental contexts: (i) short-range, (ii) long-range and (iii) chromosome-wide LD. The next section highlights the contribution of this paper and gives directions for future works. In the last section, Material and Methods, we explain the forest of hierarchical latent class models and its biological interpretation. Then, we present the new version of CFHLC able to learn FHLCMs from genome-wide data by constraining dependence scope using physical locations of SNPs. We also describe the new multilocus LD measure based on FHLCMs. Finally, we briefly explain the graph drawing- and display-methods used to visualize the FHLCMs.

Results

Short-Range Linkage Disequilibrium

We illustrate the visualization of short-range LD using the well-known Daly et al. dataset [12] available at http://www-genome.wi.mit.edu/humgen/IBD5/index.html. This dataset provides a good example of complex LD patterns with multiple degrees of LD. It consists of 129 trios, each composed of two parents and one child. For each individual, 103 SNPs are genotyped in the 5q31 region and cover 617 kb.

Our FHLCM-based method is compared with: firstly two popular approaches - the triangular heat map (THM) of $D'/\text{LOD}$ using Haplovew v4.2 (www.broad.mit.edu/mpg/) and the fine-scale recombination rates inferred with PHASE v2.1 (http://stephenslab.uchicago.edu/) - and secondly the most advanced method, the textile plot (http://www.stat.math.keio.ac.jp/). Results are presented in Figure 1. In spite of the fact that these methods differently tackle LD visualization, common trends emerge: most SNPs are divided into blocks which are common between the different methods (see dotted lines). In this sequence, Haplovew inferred 11 LD blocks, which are underlined in black color in the THM (see Figure 1a). Besides, we observe many dependences between blocks, most notably in the large central area SNP26–SNP76 (see bottom section of Figure 1a), between blocks 5 and 6. The boundary between blocks 5 and 6 is not plain. The THM also depicts strong dependences between non-contiguous SNPs, for instance, between SNPs 26 and 28 of block 5 and between SNPs of blocks 7, 8, 9 and 10. The recombination rate plot (RRP) indicates four recombination hotspots at positions SNP14–SNP15, SNP24–SNP25, SNP76–SNP77 and SNP91–SNP92, showing values beyond 10 cM/Mb (see Figure 1b). These recombination hotspots define 4 large blocks which are partly in adequacy with those obtained with the THM, as shown by the dotted lines.

In the textile plot, the greater the dispersion of the genotypes between one homozygote and the other on the vertical axis, the more likely the SNP is in LD with all other SNPs (see Figure 1c). Using the textile plot, similar results are observed compared to the THM and the RRP. Dispersions of the genotypes are high inside LD blocks and low at boundary regions. Beside the LD block view, we can distinguish the absolute LD ($r^2=1$ and $D'=1$), as observed between SNP1 and SNP2, from the complete LD ($r^2<1$ and $D'=1$), as observed between SNP2 and SNP3. For the former SNP pair, there are no connecting lines between a homozygote and the opposite side of a homozygote (e.g. GG to AA between SNP1 and SNP2), whereas there are connecting lines for the latter SNP pair. Furthermore, the textile plot offers another functionality absent in the THM and the RRP. The textile plot allows to visualize the frequencies of multilocus genotypes: the thicker the segment connecting two elementary genotypes, the higher the frequency for the corresponding two-locus genotype. For example, we observe the most frequent multilocus genotypes at the bottom section of the textile plot.

The FHLCM graph provides another view of LD (see Figure 1d). This is very similar to the THM because the method also focuses on variable dependences. In the graph, leaf nodes are SNPs (blue nodes), while the other nodes are LVs (red nodes) capturing multilocus patterns. Note that the use of other nodes (white and green nodes) will be described in the second next paragraph. An edge between two nodes (latent or observed) represents the dependence between them. Thanks to the concept of lowest common ancestor for any pair of SNPs, it is possible to gain an insight of the pairwise LD strength. In graph theory, the lowest common ancestor (LCA) is defined between two nodes.
Visualization of Multilocus Linkage Disequilibrium

(a) Blocks: 1 2 3 4 5 6 7 8 9 10 11

(b) Recombination rates (cM/Mb)

(c) Sequence information

(d) Association

Layer 0: median($r^2$) = 0.91, median(dist) = 5.5kb
Layer 1: median($r^2$) = 0.76, median(dist) = 26kb
Layer 2: median($r^2$) = 0.51, median(dist) = 61kb
Layer 3: median($r^2$) = 0.52, median(dist) = 63kb

Zoom
Figure 2. Information of the level of SNP lowest common ancestors. Relation between the level of SNP lowest common ancestors and a) the median of \( r^2 \) values and b) the median of distances computed over the corresponding SNP pairs. For instance, the left boxplot in a) concerns all pairs of SNPs with lowest common ancestors belonging to layer 1. Over these SNP pairs, the median of \( r^2 \) values is 0.91.

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number of nodes and of the number of edges. NGE varies in function of the FHLCM structure complexity. It is comprised between \( n (\text{sn}=n, \text{se}=0) \) and \( 4n^2-3 (\text{sn}=2n-1, \text{se}=2n-2) \). Obviously, this comparison is simplistic because it does not take into account the fact that the different methods do not provide the same amount of information. Nevertheless, it clearly demonstrates that, apart from the RRP, the FHLCM graph offers the best information compactness. Most notably, the comparison of the FHLCM graph with the most similar method, the THM, indicates that information compactness is much higher in the former (linear complexity) than in the latter (quadratic complexity).

Long-Range Linkage Disequilibrium

Effects Of Natural Selection. To illustrate visualization of long-range LD due to natural selection, we have chosen the major histocompatibility complex (MHC), a large chromosomal region harboring a gene family which encodes MHC molecules. MHC molecules play an important role in the immune system and autoimmunity. Long-range LD has been reported in the MHC region [16,24]. This can be explained by selective sweeps and population history (genetic drift), but there is also evidence indicating strong influence of recombination activity. To study long-range LD, we focused on the region \([22Mb-40Mb]\) present on chromosome 6 and which surrounds the MHC. Although this region contains 14292 SNPs, we preferred to select only 358 of them in the context of an off-line demonstration. It is possible to analyze all 14292 SNPs covering the MHC region but the visualization of such a large FHLCM graph requires to navigate inside the graph using software such as Tulip. We used 234 phased genotypes coming from the HapMap phase III and relative to U.S. residents of northern and western European ancestry (CEU), available at https://mathgen.stats.ox.ac.uk/impute/impute_v1.html.

The chromosome map, the triangular heat map, the textile plot and the FHLCM graph are presented in Figure 3. The three visualization methods indicate the presence of strong LD spanning several megabases on the central area (within dotted lines), surrounded by low LD regions. In the textile plot, the large strong LD region is revealed by high dispersions of genotypes at most SNPs. In the FHLCM graph, we observe several large trees and multiple layers in the central area. Compared to the textile plot, the FHLCM graph provides additional information: its multiple layers allow to distinguish between short-range LD and long-range LD. The former is displayed through low-level LCA's while the latter is highlighted by high-level LCA's. In the FHLCM, dependences between distant SNPs are easily observed. It is not the case with the textile plot where SNPs are ordered along the chromosome. To overcome this restriction, Kumasaka et al. use a hierarchical clustering variable algorithm which rearranges SNP positions along the horizontal axis, allowing to show LD between distant SNPs. Nevertheless, the drawback is that physical ordering of SNPs is lost.

Effects Of Population Admixture. We also studied the presence of long-range LD due to population admixture. For this purpose, we chose the example of the African ancestry in Southwest USA (ASW) population from the HapMap phase III. ASW is a well-known admixed population [25]. This data sample consists of 106 phased haplotypes. We focus on chromosome 1 for which we selected 2819 regularly spaced SNPs, in the context of an off-line demonstration. A sliding window of 20 Mb has been used.

The FHLCM graph is plotted in Figure 4. The graph presents 4 layers of LVA's. LD varies from \( \text{median}(r^2) = 0.053 \) in the first layer to 0.012 in the fourth layer. In the first layer, the median distance between SNPs is around 100 kb, while it is 2.66 Mb in the last layer. As observed with CMH, dependences between distant SNPs are easily observed. We are able to localize regions showing long-range LD, such as the region at the beginning of the chromosome (circled in red).

Chromosome-Wide Linkage Disequilibrium

Chromosome-wide LD visualization can be performed by learning FHLCMs with CFHLC and by navigating through the corresponding graphs with Tulip. We illustrate the LD visualization of chromosome 1 for CEU population. The dataset consists of 234 phased genotypes and 101100 SNPs. FHLCM learning was constrained by a maximum physical distance between SNPs (or LVA's) of 100 kb. CFHLC was run on a standard PC (3.8 GHz, 3.3 GB of RAM). Only 9 hours and 1.1 GB were necessary to compute the FHLCM for the entire chromosome.

In Figure 5, the FHLCM graph is depicted. Navigation in the graph through successive zooms allows to change the resolution for the visualization. When no zooming function is active, the chromosome is simply represented by a blue line. Nevertheless, if we zoom a first time on the graph, the global structure of LD becomes apparent. In the second view, long-range LD between SNPs spaced by 50–100 kb is easily visualized. In the third display, it is possible to distinguish each LD cluster in the graph (i.e. each FHLCM subtree), to see the number of FHLCM layers and the degree of connectivity. The degree of multilocus LD is shown by the color shade of LVA's. Finally, if we zoom again, we can see the position of SNPs and the precise multilocus LD strength measure, which are written inside blue nodes (SNPs) and red nodes (LVA's), respectively.

Discussion

Our FHLCM-based method for visualization of LD spatial structure has been shown to provide a compact view of LD spatial structure in the three main contexts: short-range, long-range and chromosome-wide LD analyses. Our approach focuses on variable dependences, and thus is very similar to the THM. Beside plotting pairwise LD, the FHLCM can also show multilocus LD, which represents the most important asset. Moreover, pairwise LD degrees are hierarchically displayed by LCA levels. Compared to the textile plot, our visualization tool shows several drawbacks, but also multiple advantages. Although FHLCM graphs do not allow to distinguish between complete and absolute LD, or to show genotype frequencies, it clearly reveals long-range LD without necessitating any rearrangement of the SNP order in the sequence, such as required for the textile plot. In fact, the textile plot and the FHLCM graph are complementary approaches to study LD structure.

Future researches will focus on two main aspects. First, important information provided by FHLCMs has not been used in our visualization approach. Conditional and \( \text{a priori} \) probability distributions learned by CFHLC could provide insights of the frequencies of genotypes, and above all, of the frequencies of genotype clusters. Finally, the next step is providing the geneticist an integrated software tool equipped with a user-friendly interface, such as provided by Haploviz or the Textile Plot software, to construct FHLCMs, display them and launch off-line genetic association analyses.

Materials and Methods

The Model And Its Biological Interpretation

From now on, we will restrain the study to discrete and finite variables (either observed or latent).
FHLCMs are forests whose trees are hierarchical latent class models (HLCMs). An HLCM is defined as a tree whose leaves are observed variables while internal nodes are latent variables organized in multiple layers. An FHLCM is illustrated in Figure 6. The meaning of specific key terms is clarified in Figure 7. Most notably, the benefits of using FHLCMs rely on the ability of latent layers to account for multiple degrees of SNP dependences and to naturally deal with the fuzzy nature of LD block boundaries [26]. Moreover, FHLCMs offer a generalization of the block-like structure. Inside blocks, SNPs are necessarily contiguous. For some genomic regions, the block-like structure can be irrelevant [27]. Instead of modeling blocks, FHLCMs describe clusters for which the contiguity constraint is relaxed.

In the FHLCM, LVs bring a biological meaning for the geneticist. For instance, in the case of haplotype data analysis (phased genotypes), LVs are likely to represent the haplotype
cluster structure of LD. To a certain extent, an LV might be interpreted as the shared ancestry of the haplotypes defined by the observed variables, namely, the contemporary haplotypes of the tree rooted in the LV. Each state of an LV may represent a group of similar haplotypes. In the situation of limited ancestral recombination, similar haplotypes tend to share recent common ancestry [5]. Although this situation is not guaranteed along the genome, it is very likely for low-level LVs, since they are expected to cover very small genomic regions showing strong LD. Besides, when the latent variables capture dependences between distant SNPs (or distant groups of markers), they can be viewed as population structure or natural selection effects. Thus the interpretation of LVs depends on their level in the graph. Low-level LVs covering small genomic regions in strong LD represent haplotype shared ancestry. In contrast, high-level LVs which capture weak dependences between distant SNPs correspond to population structure or natural selection effects. This distinction between multiple levels allows an easy and interpretable view of LD for the geneticist.

**FHLCM Learning**

In the CFHLC+ algorithm, the learning is performed through an adapted agglomerative hierarchical clustering procedure: (i) at each agglomerative step, a clique partitioning method is used to identify cliques of dependent variables (i.e. LD clusters); (ii) each such clique is subsumed into an LV, through a latent class model (LCM). An LCM is an HLCM which only includes one LV. For each LCM, parameter learning using the expectation-maximization (EM) algorithm and missing data imputation through probabilistic inference (for the latent variable) are performed. Iterating these two steps yields a hierarchical structure. In other words, latent variables capture the information borne by underlying observed variables (e.g. genetic markers). In their turn, these latent variables, now playing the role of observed variables, are summarized through additional latent variables, and so on. Details about clique partitioning, LCM-based data imputation for latent variables and LCM-based parameter learning of the hierarchical structure are described in the following.

**Clique Partitioning Algorithm.** A clique partition is a set of non-overlapping cliques of variables (cliques of variables can be seen as clusters of variables). The set of variables constituting the clique is likely to be subsumed into an LV. Overlapping clusters are proscribed since, in an HLCM, two latent nodes cannot share the same child. We applied clique partitioning to the complete graph of pairwise dependences. We used CAST [28], a clique partitioning method, and pairwise mutual information as a measure of pairwise dependence.

To deal with genome-scale data, a simple idea is implemented: pairwise dependences are only computed between variables (SNPs or LVs) which are separated by a maximum physical distance on the chromosome. Unlike SNPs, LVs do not have a physical location on the chromosome. To tackle this issue, for an LV, the average of the subsumed SNPs’ locations is used. The physical constraint leads to calculate a sparse matrix of pairwise dependences, where only computed values are stored. The clique partitioning algorithm CAST has been reimplemented to handle large sparse matrices.

LD modeling is constrained by the physical positions of the SNPs along the chromosome. This constraint, less drastic than genome splitting (previously proposed in Ref. [22,23]), actually corresponds to a sliding window approach. Fixing the sliding window size sufficiently large (0.1–5 Mb) represents a reasonable strategy to capture long-range LD in the GWAS context.

**LCM-Based Data Imputation For Latent Variables.** Locally, the data imputation is achieved relying on one of the simplest Bayesian networks: we consider the LCM rooted in the latent variable and whose leaves are the variables in the clique. Then, parameter learning yields the marginal distribution of the latent variable and the conditional distributions of the child variables. Parameter learning is implemented through the expectation-maximization (EM) algorithm. Subsequently, given an individual \( \ell \) and the vector of its values \( x' = (x'_1, \ldots, x'_\ell) \) corresponding to the vector of the variables in the clique \( \{X_1, \ldots, X_\ell\} \), a value \( c \) is assigned to the latent variable \( H \) through (linear) probabilistic inference:

\[
P(H = c | x') = \frac{\prod_{\ell=1}^{\ell} P(x'_\ell | H = c) \, P(H = c)}{\sum_{c=1}^{k} \prod_{\ell=1}^{\ell} P(x'_\ell | H = c) \, P(H = c)},
\]

with \( k \) the number of classes of LV \( H \). Therefore, throughout the bottom-up procedure, after the current step’s completion, the newly created latent variables will, in their turn, play the role of observed variables to seed the next step.

**LCM-Based Parameter Learning.** The role of the aforementioned local LCMs is central to the FHLCM learning algorithm. Not only do they allow data imputation for the corresponding LVs but, in parallel with the structure growing, the FHLCM parameters are also learned as follows: at \( r^{th} \) step, in addition to previously created LVs and initial OV’s not already included in the hierarchy, all LVs created at \( r-1^{st} \) step play the role of OV’s. For any such former variable shown to be a leaf node in an LCM (corresponding to a clique), the current marginal distribution is replaced with the conditional distribution learned in
the LCM. Thus, during the bottom-up construction of the FHLCM, marginal distributions are successively replaced with conditional distributions.

Implementation. CFHLC+ can process both phased and unphased genetic data. Our algorithm has been developed in C++, relying on the ProBT library dedicated to Bayesian networks (http://bayesian-programming.org). CFHLC+ is available for Windows 32 bits at https://sites.google.com/site/raphaelmouradeng/home/programs.

Multilocus Linkage Disequilibrium

Since FHLCMs represent multilocus LD, it is possible to compute a multilocus LD value from the joint probability distribution. Perhaps most interestingly, multilocus LD can be calculated for each FHLCM subtree (i.e., for each LD cluster).

Total correlation is a generalization of mutual information for multiple variables [29]. It quantifies the redundancy or dependence among a set of \( n \) random variables \( X = \{X_1, \ldots, X_n\} \). It is defined as the Kullback-Leibler divergence [30] between the joint distribution \( P(X) \) and the independent distribution \( P(X_1) \cdots P(X_n) \):

\[
C(X) = D_{KL}(P(X) || P(X_1) \cdots P(X_n)).
\]

To assess multilocus LD, total correlation over SNPs is a relevant measure to evaluate the difference between the distribution assuming linkage disequilibrium (joint distribution) and the distribution assuming linkage equilibrium (independent distribution).

Figure 5. Chromosome-wide linkage disequilibrium visualization of chromosome 1. Navigation through zooming inside the FHLCM graph. Positions of SNPs are displayed inside blue nodes. The multilocus LD strengths relative to the subtrees rooted in the latent variables can be viewed inside red nodes.

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Total correlation is reduced to the simpler difference of entropies:

\[ C(X) = \sum_{i=1}^{n} H(X_i) - H(X), \]

where the first term is the sum of entropies of individual variables and the last term is the entropy of the joint distribution of variables. The entropy \( H \) is a proper measure to assess disorder in a system \( S \) and is defined as:

\[ H(S) = -\sum_s p_s \log p_s, \]

where \( p_s \) denotes the probability of each state \( s \) in the system \( S \), and the sum includes all possible states.

Based on total correlation, we designed a multilocus LD measure relying on the learned FHLCMs. First, we need to introduce how to compute the joint probability distribution (JPD) in general Bayesian networks (BNs). Let us consider a BN modeling a set of nodes \( X = \{X_1, \ldots, X_n\} \), representing \( n \) random variables. The JPD is calculated using the recursive factorization formula:

\[ P(X) = \prod_{i=1}^{n} P(X_i|Pa(X_i)), \]

where \( Pa(X_i) \) denotes the parents of node \( X_i \), and \( P(X_i|Pa(X_i)) \) is the conditional probability distribution of \( X_i \) knowing \( Pa(X_i) \).

Now we describe the computation of the multilocus LD measure in an FHLCM subtree. In the following, we only consider subtrees composed of an LV and of all its descendants, which actually corresponds to HLCMs (e.g., the subtree rooted in node 18 in Figure 7). Let us take for example an FHLCM subtree defined on a set of \( n \) observed and latent nodes \( X = \{X_1, \ldots, X_n\} \). \( X \) is composed of a set of \( m \) observed nodes \( O = \{O_1, \ldots, O_m\} \) and a set of \( p \) latent nodes \( H = \{H_1, \ldots, H_p\} \). \( H \) can also be divided into two subsets \( N \) and \( R \); in the former, \( N = \{N_1, \ldots, N_q\} \), there are \( q \) non-root nodes, whereas in the latter, \( R = \{R_{q+1}, \ldots, R_p\} \), there are \( p-q \) root nodes.

To compute total correlation, we replace the entropy \( H(X) \), assuming linkage disequilibrium, by the entropy of the joint distribution modeled by the FHLCM subtree which is the following:
Using the mutual information formula, Equation 1 is reformulated:

\[
H(O,N,R) = \sum_{j=1}^{m} H(O_j|Pa(O_j)) + \sum_{k=1}^{q} H(N_k|Pa(N_k)) + \sum_{l=q+1}^{p} H(R_l).
\]

Thus, the total correlation \( C(X) \) writes as:

\[
C(X) = \sum_{j=1}^{m} (H(O_j) - H(O_j|Pa(O_j))) - \sum_{k=1}^{q} (H(N_k) - I(N_k,Pa(N_k))) - \sum_{l=q+1}^{p} H(R_l) = m \sum_{j=1}^{m} I(O_j,Pa(O_j)) + q \sum_{k=1}^{q} I(N_k,Pa(N_k)) - \sum_{k=1}^{q} H(N_k) - \sum_{l=q+1}^{p} H(R_l) + \sum_{k=1}^{m} \sum_{l=q+1}^{p} \sum_{j=1}^{m} H(R_l|O_j).
\]

We observe that the total correlation \( C(X) \) is composed of two terms: the former is often used to evaluate the fitness of a BN tree, or forest, without LVs (i.e., a score), whereas the latter can be seen as a penalization term specific to latent models. This penalization term is the sum of LV entropies, thus allowing to take into account the complexity increase due to the incorporation of LVs in the model. The entropy of an LV increases with both the number of its classes (i.e., states) and the uniformity of its distribution.

Finally, we scale \( C(X) \):

\[
C'(X) = \frac{\sum_{i=1}^{m} I(X_i,Pa(X_i)) - \sum_{j=1}^{m} H(O_j)}{m \sum_{k=1}^{q} H(O_k)}.
\]

In rare situations, \( C'(X) \) can be slightly below 0 (due to the penalization term). In this case, its value is set to 0. Efficient computations of \( C'(X) \) can be done starting from the bottom layer and ending at the top layer of the FHLCM. The generalization of the computation of the \( C'(X) \) measure to a whole forest is straightforward.

The interpretation of normalized total correlation \( C'(X) \) is similar to that of the \( R^2 \) coefficient, which is used for pairwise LD. \( C'(X)=0 \) means perfect equilibrium among SNPs, whereas \( C'(X) \) means that SNPs are in absolute LD (the SNPs provide exactly the same information). Note that because of the penalization term, the value of \( C'(X) \) cannot be equal to 1. Values between 0 and 1 correspond to different degrees of multilocus LD.

Graph Drawing And Visualization

Graph drawing (i.e., node placement) and visualization (i.e., display) of FHLCMs represent an important step. For this purpose, we propose a simple method which offers a clear and interpretable view of LD spatial structure. Thanks to the hierarchical nature of FHLCMs, it is possible to implement an easy and intuitive drawing: nodes are placed along the chromosome, and layer by layer. SNPs are placed along the chromosome using their physical order on the sequence. LVs are placed using physical orders computed by averaging over the orders of the subsumed SNPs. Each layer is positioned along a line parallel to the chromosome.

Regarding graph visualization, only a few software programs have been developed to handle large graphs, such as required for genome-wide LD modeling using FHLCMs. Among others, the software Tulip (http://tulip.labri.fr/TulipDrupal/) is a user-friendly tool able to deal with about one million nodes. Together with the navigation in such large-scale graphs, including zooming in narrower and narrower regions, Tulip allows the extraction of subgraphs and the enhancement of the results thus obtained by filtering. To visualize multilocus LD for each FHLCM subtree, we propose to shade the LV node subsuming the subtree, proportionally to the LD strength (\( C'(X) \)). The precise value of LD is also displayed inside this LV. Association information can also be visualized. For this purpose, the association signal is represented by an edge linking a SNP node to an additional node. The length of the edge is a linear function of the \(-\log_{10}(p-value)\) value, computed between the SNP and the phenotype. The additional node is green when the association is significant and white otherwise.

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Author Contributions

Conceived and designed the experiments: RM CS. Performed the experiments: RM. Analyzed the data: RM. Contributed reagents/materials/analysis tools: RM. Wrote the paper: RM CS PL CD.

References

1. Pritchard JK, Przeworski M (2004) Linkage disequilibrium in Humans: models and data. The American Journal of Human Genetics 69: 1–14.
2. Mc Vean G (2007) Handbook of statistical genetics, Wiley Interscience, volume 2, chapter Linkage disequilibrium, recombination and selection. 3rd edition. pp 909–944.
3. The International HapMap Consortium (2007) A second generation human haplotype map of over 3.1 million SNPs. Nature 449: 851–861.
4. Robbins MD, Sinn SC, Yang W, Deynne AV, van der Knaap E, et al. (2011) Mapping and linkage disequilibrium analysis with a genome-wide collection of SNPs that detect polymorphism in cultivated tomato. Journal of Experimental Botany 62: 1831–1945.
5. Morris AP, Cardon LR (2007) Handbook of statistical genetics, Wiley Interscience, volume 2, chapter Whole genome association. 3rd edition. pp 1238–1263.
6. Dina C, Meyre D, Gallina S, Durand E, Korner A, et al. (2007) Variation in FTO contributes to childhood obesity and severe adult obesity. Nature Genetics 39: 724–726.
7. Stephens M (2007) Handbook of statistical genetics, Wiley Interscience, volume 2, chapter Inference under the coalescent. 3rd edition. pp 478–908.
8. David J, Hand HM, Smyth P (2001) Principles of data mining. Cambridge, MA, USA: MIT Press. 578 p.
9. Simoff S, Bilen M, Maziejka A, eds. (2008) Visual data mining: theory, techniques and tools for visual analytics. Springer. 467 p.
10. Annage F, Dopazo J, eds. (2005) Data analysis and visualization in genomics and proteomics. Wiley-Blackwell. 284 p.
11. The 1000 Genomes Project Consortium (2010) A map of human genome variation from population scale sequencing. Nature 467: 1061–1073.
12. Daly MJ, Rioux JD, Schaffner SF, Hinds DA, Lander ES (2001) High-resolution haplotype structure in the human genome. Nature Genetics 29: 229–232.
13. Barrett JC, Fry B, Maller J, Daly MJ (2005) Haploview: analysis and visualization of LD and haplotype maps. Bioinformatics 21: 263–265.
14. Stephens M, Scheet P (2005) Accounting for decay of linkage disequilibrium in haplotype inference and missing-data imputation. The American Journal of Human Genetics 76: 449–462.
15. Fry B (2004) Computational information design. Ph.D. thesis, Massachusetts Institute of Technology.
16. Kumasaka N, Nakamura Y, Kamatani N (2010) The textile plot: a new linkage disequilibrium display of multiple single nucleotide polymorphism genotype data. PLoS ONE 5: e10207.
17. Koller D, Friedman N (2009) Probabilistic graphical models: principles and techniques (adaptive computation and machine learning). The MIT Press. 1208 p.
18. Thomas A (2009) Estimation of graphical models whose conditional independence graphs are interval graphs and its application to modelling linkage disequilibrium. Computational Statistics & Data Analysis 53: 1818–1829.
19. Thomas A (2009) A method and program for estimating graphical models for linkage disequilibrium that scale linearly with the number of loci, and their application to gene drop simulation. Bioinformatics 25: 1287–1292.
20. Mourad R, Sinoquet C, Leray P (2010) Learning hierarchical Bayesian networks for genotype-wide association studies. In: 19th International Conference on Computational Statistics (COMPSTAT). pp 549–556.
21. Mourad R, Sinoquet C, Leray P (2011) A hierarchical Bayesian network approach for linkage disequilibrium modeling and data-dimensionality reduction prior to genotype-wide association studies. BMC Bioinformatics 12: 16.
22. Sinoquet C, Leray P, Mourad R (2009) Inference of unexpected genetic relatedness among individuals. In: Proceedings of the 2009 IEEE International Conference on Bioinformatics and Biomedicine (BIBM). pp 524–529.
23. Mourad R, Sinoquet C, Leray P (2011) A hierarchical Bayesian network approach for linkage disequilibrium modeling and data-dimensionality reduction prior to genome-wide association studies. BMC Bioinformatics 12: 16.
24. Sato T, Kato S, Aida Y, Nakamura Y, Koide R, et al. (2003) A high-resolution linkage disequilibrium map of the human major histocompatibility complex and first generation of tag single-nucleotide polymorphisms. The American Journal of Human Genetics 76: 634–646.
25. Pemberton T, Wang C, Li JZ, Rosenberg NA (2010) Inference of unexpected genetic relatedness among individuals in HapMap phase III. The American Journal of Human Genetics 87: 457–464.
26. Anderson EC, November J (2003) Finding haplotype block boundaries by using the maximum description length principle. The American Journal of Human Genetics 73: 336–334.
27. Phillips MS, Lawrence R, Sachidanandam R, Morris AP, Balding DJ, et al. (2005) Chromosome-wide distribution of haplotype blocks and the role of recombination hot spots. Nature Genetics 33: 382–387.
28. Ben-Dor A, Shamir R, Yakhini Z (1999) Clustering gene expression patterns. In: Proceedings of the Third Annual International Conference on Research in Computational Molecular Biology (RECOMB). pp 31–42.
29. Watanabe S (1960) Information theoretical analysis of multivariate correlation. IBM Journal of Research and Development 4: 682.
30. Kullback S (1959) Information theory and statistics. John Wiley and Sons. 395 p.