Graph-based Characterisations of Cell Types and Functionally Related Modules in Promoter Capture Hi-C Data

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Keywords: Hi-C Networks, Cell Type Specificity, Graph-Based Metrics, Graph Topology.

Abstract: Current technologies, noteworthy Hi-C, for chromosome conformation capture allow to understand a broad spectrum of functional interactions between genome elements. Although significant progress has been made, there are still a lot of open questions regarding best approaches for analysis of Hi-C data to identify biologically significant features. In this paper we approach this problem by focusing strictly on the topological properties of Hi-C interaction graphs. Graph topological properties were analysed from the perspective of two research questions: 1) are topological properties alone able to distinguish between different cell types and assign biologically meaningful distances between them; 2) what is a typical structure of Hi-C interaction graphs and can we assign a biological significance to structural elements or features? The analysis was applied to a set of Hi-C interactions in 17 human haematopoietic cell types. Promising results have been obtained at answering both questions. Firstly, we propose a concrete set Base11 of 11 topology-based metrics that provide good discriminatory power between cell types. Secondly, we have explored the topological features of connected components of Hi-C interaction graphs and demonstrate that such components tend to be well conserved within particular cell type subgroups and can be well associated with known biological processes.

1 INTRODUCTION

The spatial organization of a genome inside of a living cell’s nucleus has long been appreciated as an important determinant of genomic function. One of the most important methods for studying this organization has been chromosome conformation capture or 3C, in which closely associated genomic fragments are cross-linked via formaldehyde treatment and then purified, permitting closer study (Dekker et al., 2002). Since its inception at the start of the previous decade, the original 3C protocol has been adapted and combined with other methods, including chromatin immunoprecipitation and next-generation sequencing to assist in broadening the scope and depth of individual experiments. Particularly noteworthy among these is Hi-C, which utilises a biotin-streptavidin purification method in combination with next-generation sequencing technology to potentially create a complete, unbiased genomic library of chromatin interactions (Lieberman-Aiden et al., 2009; Belton et al., 2012). However, in practice Hi-C produces highly complex datasets that do not lend themselves easily to the study of genomic contacts below a resolution of 1 Mb, and to simplify such studies an additional refinement, capture Hi-C (cHi-C), was invented. In cHi-C the complex Hi-C library is subjected to an additional sequence capture step that obtains a subset of interactions that occur with a preselected subset of genomic regions known as ‘bait fragments’, simplifying the resulting data sufficiently to allow for statistically significant results at a high genomic resolution (Dryden et al., 2014; Mifsud et al., 2015). Capture Hi-C has subsequently been used in other studies for varying elements of interest including gene promoters (Mifsud et al., 2015; Javieier et al., 2016), autoimmune disease-associated genomic variants (Martin et al., 2015), colorectal cancer risk loci (Jäger et al., 2015), cardiovascular disease risk loci (Montefiori et al., 2018) and more.

The basic analysis of Hi-C datasets is largely established and well-documented (Lajoie et al., 2015). However, the large amount of comprehensive data obtained and the high cost of exhaustive and representative experiments utilising these methods mean that it is important to consider how currently published
datasets could best be mined to answer additional biological questions. Several methods exist by which the resolution of a Hi-C dataset can be theoretically improved through more sophisticated interaction calling algorithms. These include several algorithms with broadly comparable performances for general Hi-C data (Forcato et al., 2017) and also more specific tools such as the CHICAGO analysis pipeline meant specifically for χHi-C datasets (Cairns et al., 2016), and these algorithms can involve techniques ranging from modelling technical biases (Ay et al., 2014) to employing a deep convolutional neural network (Zhang et al., 2018). Another set of approaches that are distinct from these are clustering methods where available 3C and Hi-C data are analysed in the interest of discovering functionally related modules of genes and regulatory elements. These include the use of δ-teams models for Hi-C data to identify both known and putative gene clusters (Schulz et al., 2018), hard- and soft-clustering algorithms that could theoretically assist in the interpretation of combined metagenomic sequencing and 3C data (DeMaere and Darling, 2016), and also spectral clustering-based methods such as the Arboretum-Hi-C framework that was found to be useful in identifying chromatin features in mammalian Hi-C datasets at several levels of organization and pointed to the potential utility of graph-based clustering in both analysing and comparing Hi-C datasets in general (Siahpirani et al., 2016).

Although the advantages of analysing Hi-C interaction data in graph-related terms are already well accepted – e.g. approaches by (Siahpirani et al., 2016) or (Cairns et al., 2016) explicitly discuss graph-based formalisms and their methods have been successfully applied when analysing new data sets (Javierre et al., 2016), only the properties of interaction matrices (which can be considered as weighted graphs) are mainly taken into account as well as some additional data (e.g. interaction segment distances and associations with known gene regulations). Analysis of topological features of these interaction graphs and their biological significance, however, remain largely unexplored. (Although some studies explicitly mention 'topological features', these are very limited and usually more related to the topology of chromosomes rather than interaction graphs – e.g. in (Wang et al., 2013) these are interpreted as the distributions of interaction endpoints on chromosomes).

In this paper we focus explicitly only on the topological properties of Hi-C interaction graphs. These properties were identified and analysed from the perspective of two research questions: 1) are topological properties of Hi-C interaction graphs alone able to distinguish between different cell types and assign biologically meaningful distances between them; 2) what is a typical structure of Hi-C interaction graphs and can we assign some biological significance to structural elements or features of these graphs?

2 DATASET USED FOR THIS STUDY

For this study we use a dataset of long-range interactions between promoters and other regulatory elements that was generated by The Babraham Institute and University of Cambridge (Javierre et al., 2016). The data comprise interactions that were determined by promoter capture Hi-C in 17 human primary haematopoietic cell types. The measurements have identified interaction regions of 31253 promoters across all chromosomes; from these high-confidence PCHi-C interactions have been selected using CHICAGO pipeline (Cairns et al., 2016) and selecting interactions with score 5 or more. These data are available from the Open Science Framework website (Javierre et al., 2016). This data set is still largely unique because it contains genome-wide data covering a representative subset of the entire haematopoietic lineage collected using a unified protocol.

From these data we have constructed directed graphs separately for each chromosome (few interchromosome interactions were rejected and chromosome Y was not considered due to a very few interactions) and for each cell type. The vertices of graphs are chromosome segments that correspond either to promoters (‘baits’) or the detected interaction regions (‘other ends’); the edges correspond to interactions and are directed from ‘baits’ to ‘other ends’. Unlike most other analyses we only consider the topology of interaction graphs, without assigning concrete sets of genes to the vertices. In total we obtained 23 × 17 chromosome and cell type-specific graphs, each of which can be considered as a subgraph of the ‘complete interaction network’ having a total number of vertices 251209 (with ranges between 2904 and 23079 per chromosome) and 723165 edges. Similarly, each of these graphs can be considered as a subgraph of the ‘interaction network’ of one of the chromosomes, which is specific for a given cell type.

In a number of computational tests the graphs were further modified by varying CHICAGO scores between thresholds for edge inclusion between 3 and 8. Although no comprehensive analysis was done, in general such variations had a limited impact on the stability of the results.
3 EXPLAINING DIFFERENCES BETWEEN CELL TYPES WITH GRAPH-BASED METRICS

In this section we analyse the possibility to distinguish between cell types using sets of metrics that are based on graph topological structure and propose a concrete set Base11 of metrics having the largest discriminatory power. The approach is somewhat similar to graphlet-based methods (Yaveroglu et al., 2015) that have been successfully used for analysis of different types of biological networks and also in our previous work on protein structure analysis (Viksna et al., 2004; Celms et al., 2018). The graphlet methods rely on network sampling for the presence of a predefined set of small subgraphs, for protein structure analysis the graphs were derived from typical common subgraphs.

Neither of these approaches, however, seem to be directly applicable to Hi-C interaction graphs due to lack of well defined common topological substructures (partially because the presence of edges depends on weight thresholds, and neither of the aforementioned methods are well suited to weighted graphs). Whilst features that are explicitly based on graph metrics sometimes have been used to analyse certain types of biomolecular interactions (e.g. in (Quadrini and Emanuela, 2018) for RNA structure analysis), such metrics, however, usually are more related to vertex or edge weights or labels rather than to graph topological properties.

In our set of proposed metrics we have tried to include more global graph properties, which are easy to compute and also remain comparatively stable under changing weight thresholds (at least for this particular dataset of Hi-C interaction graphs).

3.1 Graph-based Metrics

The initial set Base57 contains 57 different metrics characterising graph topology. The selection of these metrics was based on manual exploration of graphs, however, short of including all the possible counts of different small subgraphs (from which only cliques and cycles of length 2 are considered), they cover most of usually considered graph topological properties that can be computed efficiently. The set consists of the following metrics: $CCnE$ – a number of connected components with $n$ edges, $CCnV$ – a number of connected components with $n$ vertices, $BCnE$ – a number of vertex bi-connected components with $n$ edges, $BCnV$ – a number of vertex bi-connected components with $n$ vertices, $SCnE$ – a number of strongly connected components with $n$ edges, $SCnV$ – a number of strongly connected components with $n$ vertices, $BCmaxE$, $BCmaxV$, $SCmaxE$, $SCmaxV$, $BCaverE$, $BCaverV$, $SCaverE$, $SCaverV$ denote maximal and average (for cell types) numbers of vertices in connected components, and $CCmaxE$, $CCmaxV$ denote maximal and average (for cell types) numbers of vertices and edges in connected components and bi-connected and strongly connected components. Finally, $CLmaxV$ and $CLaverV$ denote maximal and average clique sizes, and $antiparallelEdges$ is number of cycles of length 2.

Apart from these additional 4 metrics were considered that are closely related to the sizes of graphs: $V$ – the number of vertices, $E$ – the number of edges, $E9$ – the number of edges, which are not shared by 9 or more (i.e. by no more than 50% of all) cell types and $E17$ – the number of edges, which are not shared by all 17 cell types.

To minimise the effect of the measure of cell type distances being based simply on different numbers of particular features (which very likely will be strongly correlated to the difference in sizes of interaction graphs) the values of metrics were normalised, i.e., for a metric $m$ its value $m(i, j)$ for cell types $i$ and $j$ was defined as $m(i, j) = m_i / \sqrt{m_i \times m_j}$, where $m_i$ and $m_j$ are the numbers of features in graphs of cell types $i$ and $j$ and $m_{ij}$ is the number of features in the largest common subgraph of these graphs. Such values $m(i, j)$ were computed for all 61 metrics and all 136 distinct tissue pairs $i$ and $j$.

One clearly expects a high correlation between the values of metrics $V$, $E$, $E9$ and $E17$; however, there are also high correlations between certain pairs from Base57 set, in particular, between numbers of vertices and edges in specific types of components. Only one of the metrics was kept for pairs with correlations of 0.93 and above (after which there is a slight drop, although the main reason for choosing this threshold was very clear correlations between metrics measuring numbers of vertices and numbers of edges), leading to the removal of $CCnV$, $BCnV$, $SCnV$ for all values of $n$ and also the removal of $CCmaxV$, $CCaverV$, $BCmaxV$, $BCaverV$ as well as $CLmaxV$ (strongly correlating with $CLmaxV$). For the further studies the remaining subset Base34 of 34 metrics from Base57 was chosen.

3.2 Cell Type Distances

While the molecular mechanisms underlying haematopoiesis in humans are incompletely understood, the general pattern of haematopoietic stem...
cell differentiation into erythroid, megakaryocytic, myeloid and lymphoid lineages is generally agreed upon and commonly represented in haematopoietic trees (e.g. Robb et al. 2007). Although hierarchical classification of blood cell types from varying lineages is occasionally performed, e.g. on the basis of ribosomal protein expression (Guimaraes and Zavolan, 2016), it is usually done with the intent of validating results by examining their consistency with well-known conceptions of haematopoiesis. The situation is similar with Hi-C data from (Javierre et al., 2016), however, this study assigns easily quantifiable distances, and, since we are working with the same Hi-C dataset, seems the most appropriate for comparison.

It should be emphasised that our goal is not to replicate the same results using slightly different methods. Our approach is distinct from (Javierre et al., 2016) in that we analyse the interactions not as a weight matrix, but in graph-topology specific terms. The question, therefore, is whether there exists a set of specific topological properties that can be utilized to distinguish between different forms of chromatin interactions captured in Hi-C data, and whether these properties are potentially useful in describing and predicting coordinated genomic processes.

On basis of these data we consider two distance measures: \( D_{cont} \) ranging from 0.00 to 1.00 that are proportional to distances from (Javierre et al., 2016) and a binarised version \( D_b \) with values 0 and 1 applying cut-off threshold 0.50 (see Figure 1). We also consider two distances based only on biologically well-accepted cell type similarities: \( D_t \) that is equal to 0 for cell types within the same (lymphoid and myeloid) subgroups and equal to 1 for cells from different subgroups, and \( D_4 \) based on their respective positions in the haematopoietic tree and (in the case of endothelial precursors \( EP \)) functional similarities. This separates cells into 5 subgroups: \[ G_1 = \{MK, Ery\}, \quad G_{2,1} = \{Mac0, Mac1, Mac2\}, \quad G_{2,2} = \{EP, Mon, Neu\}, \] \[ G_{3,1} = \{aCD4, naCD4, tCD4, nCD4, nCD8, tCD8, FoetT\} \] and \( G_{3,2} = \{nB, tB\} \) and assigns distance \( D_4 \) equal to 0.33 between different members of the same group, distances 0.67 between pairs from groups \( G_{2,1}, G_{2,2} \) and \( G_{3,1}, G_{3,2} \) and distances 1.0 between pairs from other groups. For a pair of the same cell types \( D_4 \) is 0.

In addition we also tested 3 distances \( D_A, D_B \) and \( D_C \) to check the possibility to separate some closely related cell subgroups from the other cell types in the tree (see Figure 1).

Figure 1: Haematopoietic tree of 17 cell types. The continuous distances \( D_{cont} \) range from 0 to 1 and are proportional to distances in the tree according to the scale shown. Binary distances \( D_b \) are defined to be equal to 0 for \( D_{cont} < 0.5 \) and equal to 1 otherwise. \( D_t \) is defined according to cell subtypes – it is equal to 1 between lymphoid and myeloid types and equal to 0 between cells of the same subtype. Additionally binary distances \( D_A, D_B \) and \( D_C \) were used to test separability of subtrees ‘cut’ correspondingly at points \( A, B \) and \( C \).

3.3 Discrimination Between Different Cell Types

To estimate the discriminatory power of metrics from \( Base34 \) and of 4 ‘counting’ metrics \( V, E, E9 \) and \( E17 \) we constructed linear regression models for the prediction of cell distances on the basis of the values of these metrics. We also applied stepwise regression (using Akaike information criterion (Sakamoto et al., 1986) and its implementation in \( AIC \) function, which is available in language \( R \) core library) to select the most discriminatory metrics from \( Base34 \). As statistically significant were identified 11 metrics: \( CC8+E, antiparallelEdges, CCaverE, CCmaxE, BCaverE, SCmaxE, CL3V, CLAv, CL5V, SC4E, BC8+E \) with the first 7 being significant for all four distances \( D_{cont}, D_b, D_t \) and \( D_4 \) (see Figures 2 and 3). The set of these 11 metrics is called \( Base11 \).

Some interesting features of this set are the fact that sizes of connected components have comparatively high significance, high significance has also the number of components of size 8 or more (this is also partially implied by results from Section 4). The number of cliques is also important, the dominance of \( CL3V \) could be explained by the overall larger number of them. The importance of \( antiparallelEdges \) probably could be explained by its ‘counting’ character, but potentially might be related also to gene regulation, since such interactions are possible only between ‘bait’ regions, which, by definition, are associated with genes.

The correlations between predictions of regression models and the actual distances are summarised.
in Table 1. To some extent, it shows the suitability of different metrics for discrimination between cell types, although the obtained correlation values are likely influenced by over-fitting and should be treated with caution (although the results are very stable – 10 repeated bootstrapping tests with training sets containing 75% of data produced at most 2% deviations).

An interesting feature can be observed from comparing the performance of \textit{Base11} on different chromosomes (shown by heatmaps in Figure 4). The chromosomes are grouped in a number of similarity clusters, which is not the case for randomised data. The clusters, however, strongly depend on the used cell type distance (although there are few stably related pairs of chromosomes as well as few persistent outliers).

For \textit{Dcont} distance Figure 5 shows the statistical significance of \textit{Base11} metrics for which it correlates well with metrics significance for the whole chromosome set.
Figure 4: Heatmaps of similarity between pairs of chromosomes using the four metrics giving the best predictions of cell type distances $D_{cont}$ (top left), $D_{bh}$ (top right), $D_t$ (bottom left) and $D_4$ (bottom right).

Figure 5: Statistical significance of metrics for $D_{cont}$ distance and chromosome 15, which well correlate with metrics for the whole chromosome set.

Figure 6: Statistical significance of metrics for $D_{cont}$ distance and chromosomes 19 (belongs to a block of 6 correlating chromosomes), and an 'outlier' chromosome 5.

4 THE STRUCTURE OF HI-C INTERACTION GRAPHS

In this section we more closely explore the structure of graphs describing Hi-C interactions. The initial graphs (see Section 2) are defined separately for each of the 23 chromosomes (chromosome Y is omitted due to a very low number of interactions) and for each of the 17 cell types. Each of these graphs can be considered as a subgraph of a ‘complete network’ for a specific chromosome, with vertices corresponding to chromosome segments between which interactions have been measured and (for simplicity) with isolated (non-interacting) vertices removed for a graph corresponding to a specific cell type. The number of vertices for chromosome and cell type specific graphs ranges between 773 (chromosome 22) and 11118 (chromosome 1) and the number of edges on average is slightly less than the number of vertices multiplied by 2. The vertices are denoted by the middle genomic coordinate of their respective segment mapped on a specific chromosome, and the gene annotations used are from the original data.
The questions we are trying to answer here are: 1) what is a ‘typical structure’ of Hi-C interaction graph? 2) how do the structures of graphs change if we consider common subgraphs for given sets of cell types? 3) can we assign some biological significance to the structural elements or features of these graphs?

4.1 General Topological Properties of Interaction Graphs

Since the data that we have used are focused particularly on small-scale interactions, it is not unexpected that our interaction graphs separate easily into connected components. Approximately 29% of these contain just 2 vertices (isolated edges or cycles of length 2), there are comparatively few components of size 3-7 (29%) and the remaining 45% contain 8 or more vertices. Very few of them are comparatively large (up to 150 vertices), with a typical range being 10-70 vertices and an average size being 32 vertices. A sample distribution of connected component sizes is shown in Figure 7. To investigate how these components change for different cell types we further consider components of 10 or more vertices (although chosen somewhat arbitrary, this threshold seems well suited to largely reduce random ‘noise’ in graphs). The average number of such components in each chromosome and cell type-specific graph is around 50.

As an example, the following figures illustrate a connected component of chromosome 5 and cell type EP. The complete initial component is shown in Figure 9. Figure 9 shows the parts of the component that are shared by Mac0, Mac1 and Mac2 – although a number of interactions and nodes are lost the topological structure remains mostly preserved; and also additionally is shared by cell type tCD8, in which case only a few vertices and interactions remain. The component is completely absent in cell type Neu.
Figure 10: The parts of EP Hi-C interactions component shared with cell types Mac0, Mac1 and Mac2 (top) and additionally with iCD8 (bottom). There are no common interactions shared with cell type Neu.

4.2 Biological Interpretation of Hi-C Interaction Components

The observation that connected components in Hi-C interaction graphs are either largely shared by two cell types, or are present in one of them and largely absent in another strongly suggest that they have biological roles. These, however, might differ between the components and the very large number of these make comprehensive analysis practically infeasible. Still, we have performed limited analyses that confirm a strong relation of a component structure with known biological interactions and we have explored several illustrative components in more detail.

In order to ascertain whether the clusters previously obtained by linking bait-to-bait and bait-to-end interactions contained likely candidates for functionally related gene modules we first chose a sequence of related cell types from the myeloid haematopoietic lineage to examine for cell type-specific chromatin architecture. Starting from a large pool of common connected components for resting Mac0 and inflammatory macrophages Mac1, we sequentially added more cell types to the selection criteria – alternatively activated macrophages Mac2, monocytes Mon, neutrophils Neu and endothelial precursors EP – to narrow the range of clusters and establish the specificity of graph components lost at each step.

The connected components were then assessed for the proportion of nodes lost and grouped according to their degree of preservation along the course of cell type addition, from well-preserved (75-100% of nodes in common between all cell types) to entirely lost (no nodes in common between all cell types) with several groups in between (Table 2). The majority of components found tended to be highly specific, showing 0-25% retention in the final set of linked nodes and edges – however, the most specific components also tended to have the smallest number of nodes.

Table 2: Macrophage-specific graph component retention in myeloid cells and endothelial progenitors after sequential addition of cell types. \( n \) – a number of graph components, \( n_k \) – a number of graph components containing over 20 nodes initially.

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After collecting data about node retention, the largest components showing high (75-100%) or low (0-25%) retention in chromosomes of interest according to our previous graph analysis (chromosomes 5, 9, 14, 15 and 19) were analysed for enrichment with registered transcription factor protein-protein interactions, known transcription factor binding sites, co-expressed transcription factors and binding motifs with the Enrichr web tool (Chen et al., 2013;
Figure 11: A component of Hi-C interactions shared by cell types Mac0, Mac1 and Mac2, Mon, Neu and EP. The initial larger component, shown with dashed lines, is present in Mac0.

Kuleshow et al., 2016).

The analysed lists featured between 6 and 200 genes depending on component size. The analysis uncovered a variety of transcription factors that associated with different components, including an array of broadly tissue macrophage-associated factors such as STAT5, GATA6, PPARγ and MAF (Lavin et al., 2016) as well as more specific factors found primarily in monocyte-derived macrophages – JUN, JUNB, MAFK, EGR3 and others (Ramirez et al., 2017), and even lipopolysaccharide treatment-induced transcription factors including BCL3, USF1 and SREBF2 signifying inflammatory macrophage activity or interleukin-4 and 13 activated factors such as MITF (Das et al., 2018). These specific factors were primarily, but not exclusively found in components with very little retention across all cell types analysed, such as one particular component which showed a particular enrichment for STAT5 binding during analysis and was completely lost when searching for common chromatin conformations with endothelial precursors (Figure 11).

 Altogether the gene set enrichment analysis supports the hypothesis that the genes we have linked together into connected components may form functionally related modules, broadly fitting a loose model of specificity applied through examining the overlap between chromatin architectures of diverse cell types.

### 4.3 Software Availability

Web-based software components, developed by the authors, for visualisation and exploration of Hi-C interaction graphs is available at GitHub: https://github.com/IMCS-Bioinformatics/HiCGraphAnalysis. It includes the datasets of graphs used in this study, which the users can replace with their own. The current functionality is limited to the exploration of changes in graph components being shared between different subsets of cell types.
5 CONCLUSIONS

In this paper we have analysed the topological properties of Hi-C interaction graphs from two related, but somewhat different perspectives: 1) are topological properties of these graphs alone able to distinguish between different haematopoietic cell types and assign biologically meaningful distances between them; 2) what is a typical structure of Hi-C interaction graphs and can we assign some biological significance to structural elements or features of these graphs?

In general, we think that we have obtained affirmative answers to both these questions. Regarding characterisation of interactions graphs in terms of their topological properties we propose a set Base11 of 11 metrics based on graph topological features that can be used to distinguish between different cell types for 7 different distance measures that we have tested. Although the Base11 metrics are related to graph properties, which unlikely could be directly associated with concrete biological mechanisms, they seem very appropriate for characterisation of interactions that are specific for a particular cell type. If discrimination between the cell types is the main priority then Base11 can be used in combination with ‘counting metrics’ V, E, E9 and E17 to obtain correlations between the distance-defined and predicted values of up to 0.87. Although the usefulness of these ‘counting metrics’ can be anticipated since one should expect that more closely related cell types will share more common interactions, they perform much better when used in combination with Base11, and an interesting non-obvious feature is their linearity, i.e. the fact that they perform well in linear regression models.

The observed clustering of chromosomes according to regression coefficients that are assigned to different Base11 metrics is an interesting feature, in particular since such clustering is observed for different cell type distances D, but the clusters are different for different distances D. The exact reasons why such clusters of chromosomes are formed remain unclear, but a probable explanation is that they could be the result of some complex (interaction graph-specific) dependencies between Base11 metrics, or biological relations between graph components (which is not taken into account by the current topological approach).

The analysis of topology of Hi-C interaction graphs showed that they decompose in comparatively small connected components, which can be either partially shared by all cell types, or can be pronouncedly cell type-specific – largely conserved in a specific set of cell types and practically absent in others. A detailed inspection of two of such components showed that they can be well explained by biological factors/properties that are shared by some of the cell types, but not all of them.

Regarding further developments, it would be very
interesting to test our approach on another genome-wide PCHi-C interaction dataset, in order to assess both: 1) the applicability of Base11 metrics for discrimination between cell types, other than blood cells studied here; and 2) to analyse the similarity of topological structure and component behaviour of interaction graphs in order to assess how the properties observed for haematopoietic cells generalises to other cell types. Unfortunately, as far as we know, another appropriate PCHI-C dataset covering multiple cell types has yet to become available.

There is also a good potential to further extend the graph topology based approach that we have used here. It has been quite successful to show that topological properties alone could be quite informative for discrimination between different cell types and also for assigning biological meaning to specific components on interaction graphs. At the same time some useful information in the current graph representation is absent, notably, an edge in an interaction graph might represent an interaction that forms a well-defined loop on a chromosome (if the distance between interaction segments is limited and there are no intermediate interactions between them), or it can represent a long range interaction with a far less obvious biological role. We have plans to further develop the mathematical formalism for description of interaction graphs in order to incorporate and analyse such features, although additional studies are needed to determine the best way to achieve this.

ACKNOWLEDGEMENTS

The research was supported by ERDF project 1.1.1.1/16/A/135.

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