Sequence-based modeling of three-dimensional genome architecture from kilobase to chromosome scale

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To learn how genomic sequence influences multiscale three-dimensional (3D) genome architecture, this manuscript presents a sequence-based deep-learning approach, Orca, that predicts directly from sequence the 3D genome architecture from kilobase to whole-chromosome scale. Orca captures the sequence dependencies of structures including chromatin compartments and topologically associating domains, as well as diverse types of interactions from CTCF-mediated to enhancer-promoter interactions and Polycomb-mediated interactions with cell-type specificity. Orca enables various applications including predicting structural variant effects on multiscale genome organization and it recapitulated effects of experimentally studied variants at varying sizes (300 bp to 90 Mb). Moreover, Orca enables in silico virtual screens to probe the sequence basis of 3D genome organization at different scales. At the submegabase scale, it predicted specific transcription factor motifs underlying cell-type-specific genome interactions. At the compartment scale, virtual screens of sequence activities suggest a model for the sequence basis of chromatin compartments with a prominent role of transcription start sites.

Understanding how genomic sequence directs genome folding into 3D structures at all spatial scales will be instructive in interpreting how genomic sequences and genome variations are involved in various cellular processes (for example, gene expression regulation, DNA replication and DNA repair) under both normal and disease states. Such sequence dependencies are likely multifold, as there are multiple facets of 3D genome organization that appear to correspond to distinct mechanisms. Most prominently, chromatin compartments are observed typically at megabase-scale with a characteristic plaid-like interaction pattern, where compartments A and B largely correspond to expression-active and -inactive chromatin, which preferentially interact with the same compartment1. Topologically associating domains (TADs) are found at typically 100-kb to 1-Mb scale1–3 with an often nested structure.

Despite known associations with gene expression activity and specific histone marks4–8, the sequence basis of large-scale organization of chromatin compartments remains unresolved. At submegabase-scale, the formation of TADs is well known to be dependent on CTCF sequence motifs1–3, likely through a CTCF-cohesin-dependent loop extrusion mechanism10–12. However, the sequence determinants of multiple types of CTCF-independent interactions, including enhancer–promoter interactions and Polycomb-induced contacts, are less well understood, let alone predicting these interactions from sequence.

The development and improvement of high-throughput chromatin conformation capture (3C) (ref. 13)-based methods, including Hi-C and micro-C14,15, comprehensively catalog diverse types of genome interactions from kilobase to whole-chromosome scale. This provides the foundation for developing machine-learning approaches to recognize the complex sequence dependencies of genome interactions.

Learning the sequence dependencies of 3D genome structure across spatial scales will importantly provide the ability to predict the impact of new sequences. Predicting multiscale 3D genome structure from the sequence will not only enable the prediction of the impact of any sequence variant but also help to understand new sequence-based mechanisms of 3D genome organization. Deep-learning sequence models have been applied to modeling various biochemical and regulatory properties based on genomic sequences16–22. Recent works including Akita23 and DeepC24 have led to a breakthrough in deep-learning sequence-based modeling of submegabase 3D genome structure, which allows prediction of genome interactions up to 1-Mb distance from genomic sequence. However, no sequence models that predict large-scale genome organization involving sequence context beyond 1 Mb have been developed. This limits our ability to predict large structures, including chromatin compartments and local structures that depend on larger sequence context. Moreover, the lack of large-scale sequence models also limits our capability of modeling effects of large structural variants (SVs), which are among the most impactful genomic variations.

To enable modeling all scales of genome architecture measured by Hi-C-type methods, I developed Orca, a multiscale sequence modeling framework that predicts from sequence the 3D genome structure from kilobase-scale up to whole-chromosome scale, as measured by 3C data. Orca enables the prediction of diverse types of structures including TADs, chromatin A/B compartments, Polycomb-mediated interactions and promoter–enhancer interactions. Moreover, both intrachromosomal and interchromosomal interactions, from any pair of sequences in the genome, can be predicted with this approach.

Orca sequence models effectively provide an ‘in silico genome observatory’ of 3D genome architecture that uniquely enables (1) predicting the multiscale 3D genome organization effects of any genome variant of any size in high throughput, and (2) designing and performing ‘virtual genetic screen’ experiments that probe sequence-based mechanisms of multiscale genome organization. The models’ capabilities in predicting 3D genome effects of diverse SVs were extensively studied and the models were applied to generate hypotheses for the sequence-based mechanism of local genome
organization and chromatin compartment formation. The Orca sequence modeling framework can provide new opportunities for studying the interplay between sequence and multiscale 3D genome organization. The code and models can be accessed from https://github.com/jzhoulab/orca and a user-friendly webservice is available at https://orca.zhoulab.io.

Results

Sequence-based prediction of multiscale 3D genome interactions. Chromatin organization at multiple scales shows distinct characteristics and likely involves varied mechanisms, and capturing sequence dependencies across all scales from single nucleotides to the entire chromosome with deep learning is an unprecedented challenge. A multiscale deep-learning sequence modeling framework, Orca, was first developed to address this challenge.

To predict across the whole range of genomic-distance scales, a ‘zooming’-like cascading prediction mechanism was designed to enable the prediction of ultra-long-distance interactions to shorter-distance interactions with nine different resolutions (for example, 4 kb at 1-Mb distance, 8 kb at 2-Mb distance and 512 kb at 128-Mb distance). Since Hi-C-type data are typically represented through multiresolution matrices\(^{16,28}\), and longer-distance large-scale structures are typically detected based on sparser sequencing reads and thus can only be measured with a lower resolution, modeling multiscale structure at different resolutions is designed to fit these data types.

The model architecture is composed of a hierarchical multi-resolution sequence encoder and a cascading multilevel decoder. The encoder takes up to 256-Mb sequence as input and generates a series of increasingly coarse-grained sequence representations at nine resolutions from 4 to 1,024 kb. The multilevel decoders predict up to 256-Mb-distance interactions at the top level, which is larger than the longest human chromosome chr1, and down to 4-kb-resolution interactions within 1-Mb distance at the bottom level. Interchromosomal interactions are also allowed at 32–256-Mb levels by using multichromosomal input (Methods). The detailed multiscale deep-learning sequence model architecture specification is provided in Supplementary Fig. 1 and the code repository.

To enable scaling of deep-learning model training and inference to large chromosome-scale sequences, a horizontal checkpointing technique for increasing memory efficiency (Methods) was devised to allow training models even when the internal representation size far exceeds the GPU memory bound.

Orca sequence models were trained on the micro-C datasets for H1 embryonic stem cells (H1-ESCs) and human foreskin fibroblasts (HFFs), which are among the highest-resolution datasets to date\(^{11}\). The encoders and decoders are jointly trained in three stages, during which the encoder trained in the earlier stage is frozen and used in the later stage training (Methods). The final models predict from 1 to 256 Mb at nine different scales (Fig. 1a–c). Each model consists of 1-Mb, 1–32-Mb and 32–256-Mb modules that can be used together or separately to provide flexibility in applications: the 1-Mb model is the main model with high accuracy and flexibility for most applications; the 32–256-Mb model is most useful for prediction of chromosome-scale and interchromosomal interactions; and the 1-Mb model is useful for rapid screening of local genome interaction effects for a very large number of variants.

The predicted interaction matrix scores represent the log fold over distance-based background scores, where the background scores (also often referred to as the expected scores) are the average normalized contact score at the same genomic distance. On holdout test chromosomes, the model achieves 0.78–0.85 Pearson correlation with experimental observations consistently across all scales for H1-ESCs and 0.73–0.79 Pearson correlation for HFFs (Fig. 1c and Extended Data Fig. 1). Interchromosomal interactions are predicted with correlations of 0.47–0.74 (Supplementary Fig. 2, 64–256-Mb scale structural impact prediction was first demonstrated with a large 40.5-Mb inversion mutation that was found to be a potential cause of acute myeloid leukemia\(^{27,30}\), and the predictions were shown

Multiscale SV effects on 3D genome predicted from sequence. Since Orca models can accurately predict genome interactions across scales from new unseen sequences, they could be particularly useful when applied to predicting genome variation effects. Notably, because Orca allows very large sequence input (256 Mb, larger than the longest human chromosome chr1: 249 Mb), it enables predicting effects for variants of nearly any size, including very large SVs and copy number variants which are among the most impactful genome variants\(^{17,28}\). To predict genome structural impact of any variant, one can computationally reconstruct the chromosomal sequence that carries the variant, and compare the predictions against the predictions of the reference sequence. Joint effects of multiple variants on the same haplotype can also be predicted in a similar fashion.

The SV effect predictions of transposon-mediated 2-kb TAD boundary element insertions into various genome locations (2-kb insertion + 5-kb transposon), which have been measured with in situ Hi-C\(^{29}\), were first tested. The insulation score change at each insertion site was computed and compared with the predicted changes. Across 14 insertion sites, Orca attained a cosine similarity score of 0.89 for the H1-ESC model and 0.76 for the HFF model for insulation score changes (P < 1 × 10\(^{-4}\) for both H1-ESC and HFF; Methods). Moreover, Orca predictions recapitulated all three categories of insertion effects reported, including formation of new boundaries, strengthening existing boundaries and no domain-level effects (Extended Data Fig. 5 and Supplementary Data 2). Thus, the experimental Hi-C measurements are highly consistent with the Orca predictions on the genome structural effects of these insertions.

To evaluate the model’s capability in predicting the impacts of diverse types of SVs, the effects of a variety of SVs ranging from 0.3 kb to 80 Mb in size with experimentally measured genome structural impact were predicted (Supplementary Table 3, Fig. 2, Extended Data Fig. 6 and Supplementary Figs. 8–14). The multiscale structural impact prediction was first demonstrated with a large 40.5-Mb inversion mutation that was found to be a potential cause of acute myeloid leukemia\(^{27,30}\), and the predictions were shown
**Fig. 1 | Predicting multiscale 3D genome architecture from sequence.**

**a.** Schematic overview of the deep-learning model architecture for genome interaction prediction across all scales. Sequence representations at multiple resolutions are computed by a hierarchical encoder starting from the sequence in a bottom-up (high-resolution to low-resolution) order, whereas genome interaction matrices are predicted from both the corresponding levels of sequence representation and the higher-level genome interaction prediction in a top-down order (low-resolution to high-resolution). **b.** Multiscale sequence-based prediction example zooming from the whole chromosome into a position on a holdout test chromosome. Predictions from 1–256-Mb scales are compared with micro-C experimental observations. Missing values in micro-C data due to lack of coverage are shown in gray, and these regions are also indicated in the 64–256-Mb predictions because the predictions at major assembly gaps or unmappable regions are of unknown accuracy. The genome interactions are represented by the log fold over genomic-distance-based background scores for both the prediction and the experimental data. The predictions for the same regions for the HFF cell type are also shown in Extended Data Fig. 1. **c.** Scatter plot comparison of the predicted interaction scores with the micro-C measured interaction scores on the holdout test chromosomes. In each panel, 10,000 randomly subsampled scores are shown. The overall Pearson correlations across the entire test chromosomes are also annotated. Predictions for 1–32-Mb levels are from the Orca 32-Mb model and 64–256-Mb levels are from the Orca 256-Mb model.
Multiscale sequence-based prediction of SV effects on genome structure. **a.** Schematic illustration of sequence-based predictions of multiscale genome interaction effects of SVs. A large 40.5-Mb inversion variant involved in leukemia is shown as an example. Predicted effects are shown by predicted genome interaction matrices based on wild-type (WT) sequences and mutated sequences (Mut) at multiple scales. The experimentally supported effects of SVs are illustrated at the top of each panel (**a**–**c**), with relevant gene positions, major TAD boundaries (marked with the letter B) and the range of variant positions indicated (minimal variant range indicated in bold lines). Experimentally supported increase in ectopic interactions is indicated with blue dashed arcs and blue bars. The Orca genome interaction predictions are represented by the log fold over genomic-distance-based background scores. **b.** Orca predictions of multiple variants with complex phenotypic outcomes in WNT6-PAX3 region. Positions of the major genes affected by the SVs are indicated by black arrows and known enhancer regions involved are indicated by blue arrows. Ectopic interactions caused by the variants are indicated by circles. Black and gray bars on the left side indicate genomic intervals involved in the SVs pre- and post-mutation. Full multiscale prediction results for both H1-ESC and HFF cell types as well as micro-C observations in the cell types are included in Supplementary Data 3, and validation results for all SVs are summarized in Supplementary Table 3. **c.** Comparison with 4C-seq experimental data for variants predicted in b. The normalized counts from 4C-seq and log_{10} predicted interaction scores (fold over background) at the 4C-seq point-of-view are shown. The observed and predicted gains of interaction sites relevant to the phenotype are highlighted with the red dashed line box. 4C-seq, circular chromosome conformation capture sequencing; Del, deletion; Dup, duplication; Inv, inversion.
at five different levels zooming from a whole chromosome view into the EVII-proximal breakpoint (Fig. 2a and Supplementary Fig. 8; full predictions at all scales also available in Supplementary Data 3). The predictions showcase both large-scale remodeling of the chromosome organization and the breakpoint-adjacent effects on chromatin compartments and TADs, including at the finest level a gain of EVII promoter interaction with a GATA2 enhancer, which has been experimentally confirmed\(^6\).

Orca predictions were next applied to analyze a complex region where multiple deletion, inversion and duplication variants ranging from 0.9 Mb to 1.8 Mb lead to several different limb malformation phenotypes: brachydactyly, F-syndrome and polydactyly\(^6\). Orca predicts that through different structural alterations all these disease SVs cause de novo contacts between three different genes, PAX3, WNT6 and IHH, with the same enhancer region (Fig. 2b,c and Supplementary Fig. 9). These predictions are also fully consistent with previous experimental data based on circular chromosome conformation capture (4C) experiments\(^1\). These variants showcase several distinct mechanisms that create ectopic interactions predicted by the sequence models: fusion of TADs by boundary deletion creates interactions between distal positions that belonged to two different TADs (PAX3 and the enhancer); duplication creates ectopic interaction between WNT6 and the same enhancer by placing the WNT6 sequence into a new context, with similar mechanisms also observed for the Cooks syndrome duplications; inversion that spans a TAD boundary leads to changed compositions of both TADs, which results in ectopic interactions of IHH with sequence from a different TAD.

Orca was also applied to complex genomic regions where several adjacent SVs lead to distinct outcomes (Supplementary Figs. 10–14, Extended Data Fig. 6 and Supplementary Data 3). The KCNJ2-SOX9 region was first studied, where duplication variants (length 0.2–1.9 Mb) are observed to cause three distinct outcomes: sex reversal (female-to-male), Cooks syndrome (finger hereditary disorder) and no phenotype. Notably, the no phenotype duplication fully encompasses the sex reversal duplication regions. This region has been carefully studied experimentally in ref. \(^1\). The effects of both long- and short-form SVs that lead to each phenotype were predicted. Each of the variants is visualized at a selected scale to showcase their impacts in Fig. 2, and the full predictions are available in Supplementary Data 3.

Orca’s sequence-based predictions show that sex reversal duplications (0.2–1 Mb) lead to an enlarged TAD with duplicated interactions with SOX9 within the TAD (Supplementary Figs. 10–12 and Extended Data Fig. 6). The duplicated regions include an enhancer\(^1\) in both maximal and minimal duplication variants that cause sex reversal, creating de novo contact between SOX9 and the new copy of the enhancer (Supplementary Fig. 10). In contrast, the larger no phenotype duplication (1.8 Mb), which also includes the RevSex region, is predicted to leave the genome interaction patterns of SOX9 unchanged despite the duplication, because the duplication established a new TAD boundary insulating the new copy from interacting with SOX9. This explains why these duplications lead to the ‘no phenotype’ outcome (Supplementary Figs. 10 and 13). A third distinct outcome from disruptions of this region, Cooks syndrome (a finger hereditary disorder), is caused by duplications further extended to include the KCNJ2 gene (1.4–2 Mb). The model predicted that the new copy of KCNJ2 is located in a newly formed TAD due to the duplication, with KCNJ2 hijacking the genome interactions of SOX9. Similar to the no phenotype duplications, SOX9 is insulated in its original TAD and its interactions remain unaffected (Supplementary Figs. 10 and 14). Therefore, these results present models of 3D genome architecture changes that explain the phenotypes of those variants, which are fully in agreement with experimental data\(^1\). The predictions also provide extra support for the proposed structural changes by resolving ambiguity from the sequencing-based experimental results due to the two duplicated regions being indistinguishable from sequencing reads.

Overall, Orca predictions of SV effects were tested on a diverse set of variants across six studies and, remarkably, in all cases the predictions are concordant with the experimental observations with chromatin conformation capture experiments (Fig. 2, Supplementary Figs. 8–14 and Supplementary Data 3, and summarized in Supplementary Table 3). Importantly, such sequence analysis can be made in seconds and is thus scalable to millions or more variants. The accurate recapitulation of genome organization effects of these SVs shows the potential in predicting structural effects of variants without previous data on their consequences in genome 3D structural organization.

**Motifs underlie cell-type-specific local genome interactions.** The model’s capability in predicting 3D genome architecture at multiple scales directly from sequence also allows to use it as an ‘in silico genome observatory’ to probe the sequence determinants of 3D genome organization learned by the deep-learning models. This computational approach has the capability of performing ‘virtual genetic screens’ on a very large number of sequences and allows almost unlimited flexibility in sequence design. Here multiple screen strategies were designed for dissecting the sequence basis of local (1-Mb) and compartment-level (32-Mb) organization, which revealed distinct sequence dependencies.

For discovering sequence dependencies of the submegabase-scale genome structures that are exemplified by TAD, sub-TAD and promoter–enhancer interactions, a multiplexed in silico mutagenesis approach was devised to screen for sequence disruptions that lead to ‘local’ structural remodeling within 1-Mb distance (Fig. 3a). This multiplexed approach introduces multiple site-disruption mutations to the same 1-Mb sequence to speed up near-basepair-level screens, leading to 20× speed-up in this example. Moreover, each 10-bp site is disrupted in three different sequences, each with a random set of disruption sites. Leveraging the sparsity of mutations with structural impact, the site-specific effects were deconvolved using the minimum 1-Mb structural impact score (average absolute log fold interaction changes between the disrupted position and all other positions within the 1-Mb window) across three sequences sharing the same disruption site as the final score (Methods). Taking the minimum of multiple sequences each with independent random disruptions also has the advantage of filtering out the low-probability events caused only by specific mutated sequences. We show that this multiplexed approach is highly concordant with the single-mutation approach with >0.9 correlation (Extended Data Fig. 7).

With this approach, all 10-bp sequences on autosomes whose disruptions have structural impact were screened. Consistent with the central role of CTCF in TAD-level structural organization, for both H1-ESCs and HFFs, most of the 10-bp sites (>88.9%) with the strongest tier of 1-Mb structural impact score (>0.1, <0.015% of the genome) are overlapping with CTCF motifs (log-odds >10) (Fig. 3b) and >95.1% are within 200-bp distance to a CTCF motif (Supplementary Fig. 15), while only <1% are depleted of CTCF motifs (log-odds <6) within 200-bp distance (versus a genome-wide background of 64%). This suggests that the strongest impact sites are predominantly explained by CTCF. However, not all CTCF motifs are predicted to have strong structural impact (only ~1% of sites with CTCF motif log-odds >10 have structural impact score >0.1); thus, the CTCF motif is not the only determinant and the model utilizes more complex sequence dependencies to make accurate predictions.

Despite that the strongest tier of 1-Mb structural impact score sites are predominantly CTCF related, non-CTCF transcription factor motifs are highly enriched in the mid-impact score range
In contrast to the CTCF motif dependency, which is largely cell-type-invariant (Fig. 3b), very strong cell-type specificity was observed in non-CTCF motifs that are predicted to impact genome structure: H1-ESC is predicted to be most responsive to the disruption of the POU5F1::SOX2 dimer motif and POU family motifs, while HFF is highly sensitive to AP-1 (FOS::JUN) motif disruptions (Fig. 3c, Supplementary Tables 4 and 5 and Supplementary Fig. 16; the POU5F1::SOX2 motif is $48.7\times$ and 1.0x enriched in H1-ESC and HFF, and the FOSL1::JUND motif is $0.71\times$ and 167x enriched in H1-ESC and HFF, with motif log-odds >12). This cell-type selectivity is consistent with the well-known gene regulatory roles of POU5F1 and SOX2 in embryonic stem cells and AP-1 in fibroblasts. Example disruptions of single POU5F1::SOX2 or AP-1 motifs can lead to the elimination of predicted genome interactions in H1-ESC and HFF cell models (Supplementary Figs. 17 and 18). These results suggest that cell-type-specific transcription factors mediate local interactions and may also impact transcription through the spatial organization.
A sequence basis model for compartments with transcription start sites (TSSs) as drivers. As the Orca sequence models uniquely enable the prediction of compartment-level (>1-Mb) genome interactions from sequence, it presents an opportunity to probe into the sequence-based mechanisms of chromatin compartment formation. To understand the sequence dependencies of the compartment-level genome structures as learned by the model, the challenge of deconvoluting sequence effects on chromatin compartments from CTCF-cohesin-mediated mechanisms such as TAD organization needs to be first addressed. To differentiate sequence effects on chromatin compartments, a new sequence model was trained with a cohesin-depleted Hi-C dataset for HCT116 cells. As acute cohesin depletion completely eliminates TAD domains while chromatin compartments remain intact or strengthened, this sequence model learned only sequence dependencies of chromatin compartments but not CTCF-cohesin-dependent structures. The cohesin-depleted HCT116 sequence model predicts genome interactions with a Pearson correlation of 0.71 (32-Mb level, on holdout test chromosomes) and no apparent CTCF motif dependency was observed in this model, consistent with the clean elimination of TADs in this dataset (Supplementary Fig. 19).

To identify the characteristics of sequences that are sufficient for establishing the generally expression-active A or inactive B chromatin compartment according to the model, a virtual genetic screen for ectopic sequence activity in switching A/B chromatin compartment was designed with a strategy of swapping in ‘insertion’ sequences from positions of the genome to a diverse set of target positions (Fig. 4a).

The main characteristics of sequence chromatin compartment activities were here demonstrated with a screen involving a set of 2,500×12,800-bp source sequences that tiled a 32-Mb region with multiple A and B compartment regions, and 10 target insertion sites uniformly spaced in the same region (Fig. 4b). For each target site, visualizing the predicted 32-Mb structural impact score (average absolute log fold change in interactions with the insertion site within the 32-Mb window) of all source sequences against the source sequence positions generates a sequence compartment activity profile. These sequence activity profiles are clearly grouped by the compartmental context of the target sites (Fig. 4c, Extended Data Fig. 8 and Supplementary Fig. 20): compartment B target sites which detect predominantly B-to-A (B→A) compartment change and compartment A target sites which detect predominantly A-to-B (A→B) compartment changes, and sites that are near compartment boundaries which detect mixed changes. For succinctness, insertion sequences that are predicted to cause B→A and A→B changes will be referred to as having compartment A activity and compartment B activity, respectively.

Sequences with compartment A activities are sparsely distributed in predominantly compartment A regions (Fig. 4c,d; <0.07% of 400-bp sequences have >0.02 32-Mb structural impact score). In contrast, sequences with compartment B activities are widespread across all compartment B regions (Fig. 4c,d). Closer inspection revealed that sequences with strong compartment A activities mostly span TSSs (Fig. 4d,e). Similar results were recapitulated with a large-scale screen using 12,800-bp insertion sequences tiling all of the holdout chromosomes and 200 random target sites (Fig. 4f, Extended Data Fig. 8 and Supplementary Fig. 20). In contrast, the 3’ ends of genes are not enriched in compartment A activity (Supplementary Fig. 21). In addition, overlap with HCT116 chromatin states shows very strong enrichment in active TSS states but not in bivalent/poised TSS or other states (Supplementary Fig. 22). Furthermore, little TSS transcription directionality preference was observed in all 200 target positions (Supplementary Fig. 23). A model of active TSS sequences being drivers for compartment A is also in line with previous observations that linked transcription with compartment. It is worth noting that the cohesin-depleted HCT116 sequence model learned to recognize TSS sequences while training on only 3D genome data without any transcription or chromatin profile data.

The minimum length of insertion sequence that is required for chromatin compartment switching was then assessed using a series of insertion sequence sizes ranging from 200 bp to 51,200 bp (Fig. 4d). Remarkably, insertion of only 800-bp sequence is sufficient for strong A compartment activity, and even 400-bp or 200-bp sequences can have partial effects (Fig. 4d,e). Further increase of length does not significantly increase ectopic compartment A activity until the length starts to cover multiple TSSs. While sequences as short as 800 bp are sufficient for the establishment of A compartment within a native B compartment environment, these sequences are expected to induce widespread chromatin changes due to transcription activity or histone modification. For detectable A→B directional compartment change at a native A compartment environment, a minimum of 6,400 bp is needed, while 12,800 bp or longer produces more pronounced effects. Interestingly this 6–12-kb minimum length scale coincides with independent experimental measurements of minimum DNA fragment length for maintaining stable compartmentalization. Fragments of at least 10–25 kb are required for stable compartmentalization while <6-kb fragments lead to a gradual loss of genome organization.

Finally, the size of sequence patterns necessary for compartment activity was analyzed via a permutation-based approach. Specifically, the insertion screen was modified by first dividing the sequence into segments of the same size and then randomly shuffling the order of the segments before insertion, and the effects of different segment sizes were compared. Since permutation eliminates all sequence patterns larger than the specified segment size, if the sequence activity is unaffected after permutation, then such
activity only depends upon sequence patterns smaller than the segment size. Interestingly, predicted A-to-B compartment change can be achieved with the insertion of randomly permuted sequences originally with B compartment activity at essentially all segment sizes down to 2 bp (Extended Data Fig. 9). Thus, complex sequence patterns beyond mono- or dinucleotide frequencies may not be necessary for strong predicted B compartment activity (this does not preclude the possibility that some complex sequence patterns can be
sufficient for B compartment formation). B compartment activities, before and after permutations, are also correlated with high A/T, low G/C content, as well as AT dinucleotide pattern (Extended Data Fig. 9 and Supplementary Fig. 24), which is also consistent with the known enrichment of A/T in the B compartment. In contrast to B compartment activity, compartment A activity is nearly eliminated with segment size below 128 bp and is also decreased at 256 bp; thus, compartment activity likely depends upon sequence patterns of size at least 128–256 bp, which is in line with the involvement of TSS sequences. Moreover, the permuted sequences not only lose A compartment activity but also even gain weak compartment B activity (Extended Data Fig. 9). Consistently, disruptions of extended regions of compartment A sequences (for example, 1.28 Mb) by random permutations are predicted to lead to B compartment formation, while randomly permuted compartment B sequence remains B compartment (Extended Data Fig. 10).

Taken together, these results suggest a sequence-oriented model postulating that chromatin compartment A formation is driven by TSS sequences, likely through induced transcriptional activity and chromatin state changes, while compartment B requires extended sequences (>6–12 kb) without compartment A activity, has a preference to AT-rich sequences and may be the ‘default’ state established on all non-compartment A sequences.

Discussion
Orca is a sequence model framework for global prediction of 3D genome organization across spatial scales from kilobase to whole chromosomes, based on only genome sequence. It allows the prediction of genome structural impacts of any genome variant including large structural and copy number variants. Orca accurately recapitulated the structural impacts of variants that have previously been experimentally studied. With the potential of rapidly analyzing a large number of variants requiring only the sequences, it can help accelerate the study of SVs’ roles in health and disease. In addition to enabling predicting variant effects at scale, these sequence models that capture sequence dependencies of 3D genome interaction structures provide tools for probing sequence-level mechanisms of genome interactions with virtual genetic screens.

As with the multiscale spatial organization of the 3D genome, the sequence dependencies are expected to vary by scale. Sequence determinants at the scale of a single motif appear to be a combination of strong-effect CTCF motifs and medium to weak effect tissue-specific TF motifs, possibly through different mechanisms. At hundreds of basepairs length, sequences at TSSs are predicted to have activity for establishing compartment A. At 6–12 kb and above, extended stretches of B compartment sequences or even randomly scrambled sequences can establish B compartment. Recently experimentally determined minimal length of genome fragments for maintaining compartment structure is around 6–10 kb (ref. 29), which is similar to the length scale required to induce significant A > B compartment change. This may suggest this is a key length scale that is required for the underlying biophysical mechanisms of compartmentalization, possibly through phase separation.

From a sequence-based perspective, compartment A appears to be the ‘active’ compartment that requires specific sequence patterns, as widespread chromatin changes may be induced with the insertion of TSS-proximal sequences. In contrast, compartment B appears to be the ‘passive’ compartment as it requires extended sequences without compartment A activity, and compartment B structures are predicted to be robust to random permutation of sequence. Note that the notion of ‘active’ or ‘passive’ here indicates only the sequence dependency characteristics but not the molecular mechanisms, as establishment and maintenance of both compartments could involve active molecular biochemical activities. These hypotheses remain to be tested through future experiments. Moreover, further studies may extend or revise sequence dependencies of chromatin compartment that are proposed in this model, such as possible dependencies on sequences that activate or repress transcription.

There are a few limitations of this study that are worth mentioning. Even though the predictions closely recapitulate experimental observations in most cases, in some cases they still differ from observation beyond what can be explained by technical noise or alignment artifacts as shown in Supplementary Data 1. Thus, there is still space for further improvement in performance, and new sequence-based mechanisms are expected to be discovered with higher-resolution data and improved models. Secondly, machine-learning-based approaches such as Orca are expected to capture sequence pattern dependencies that recur across the genome, and therefore sequence-based mechanisms that uniquely apply to very few or even a single genomic locus may not be learned through this approach. Thirdly, sequence models may face challenges in learning the correct ‘driver’ sequence patterns when the ‘passenger’ sequence patterns are nearly perfectly correlated with the driving factor, even though the model can identify the correct driver when the correlations are less perfect and the training data are sufficiently informative. Lastly, because of the current limitation in measuring structure for highly repetitive regions from Hi-C reads, the model’s predictions in these regions were unable to rigorously assessed (the model typically predicts B compartment-like structure for these regions). Complete end-to-end assembly of the human genome and long-read sequencing techniques may allow addressing of this limitation in the future.

The Orca sequence models also provide ample new opportunities for designing sequence-based experiments to probe sequence dependencies of 3D genome organization with ‘virtual genetic screens’ beyond what was explored in this manuscript, such as finely dissecting sequences at basepair resolution for interactions at specific loci. Such analyses can be done with the models and code released here. More generally, I anticipate such deep-learning model-based approaches for in silico modeling of complex biological processes to be powerful methods to generate hypotheses for biological systems.

Online content
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Methods

Orca model architecture for multiscale 3D genome prediction. The Orca model architecture is composed of a hierarchical sequence encoder and a multilevel cascading decoder, designed to provide a ‘zooming’ series of predictions at multiple scales. The 1D-to-4D hierarchical sequence encoder transforms raw sequence input up to 256 Mb to a series of sequence representations at multiple resolutions. A series of cascading decoders each predict an interaction matrix, which represents all pairwise genome interactions within a window of varying sizes from 1 to 256 Mb at different resolutions. All predicted interaction matrices are of size 250 × 250 and all predicted scores represent log fold over distance-based background. The decoder at each level takes the sequence encoding at the corresponding resolution as input. The top-level decoder receives input from the entire sequence at the lowest resolution, and lower levels receive sequence representations at higher resolutions. For example, the 32-Mb level decoder receives 128-kb resolution sequence encoding for 32-Mb sequence and the 1-Mb level decoder receives 4-kb resolution sequence encoding for 1-Mb sequence. In addition, except for the top-level decoder, lower-level decoders also receive the prediction from the upper level as input (for example, 1-Mb level decoder receives 2-Mb level prediction, cropped to the 1-Mb region), and all multilevel decoders also receive a distance encoding matrix as input. The encoder computation starts from a bottom-up pass (high resolution to low resolution) starting from raw sequence with one-hot encoding, followed by a top-down pass to introduce longer-range information to the finer-resolution representations (Supplementary Fig. 1). The decoder computation follows a top-down order (long maximum distance to short maximum distance, low resolution to high resolution) and each lower-level decoder receives the upper-level prediction from the upper level as input. The architecture and input are described in more detail below, and the detailed architecture of models is available in Supplementary Fig. 1 and the code repository.

Both the encoder and decoders are convolutional networks with residual connections. The hierarchical sequence encoder alternates between one-dimensional (1D) residual convolution blocks and max-pooling layers. More specifically, the first section of the sequence encoder converts the one-hot sequence encoding into 4-kb-resolution sequence representations with a convolutional architecture adapted from the Seti model58 that uses a dual linear + nonlinear path design that stacks nonlinear blocks with residual connections on top of the linear blocks (Supplementary Fig. 1). The first section of the encoder contains 28 convolution layers each with 64–128 channels. With the 4-kb resolution sequence encoding as input, the upper sections of the encoder create a series of sequence encoding at 4 kb, 8 kb, …, 1,024 kb resolutions with factors of 2 with a similar residual block structure, using 4 convolution layers per resolution with 128 channels.

To predict two-dimensional (2D) interaction matrices at multiple scales, a cascading series of sequence decoders was used, each predicting a genome interaction matrix with a different length and resolution. The 2D convolution architecture consists of 2D residual convolution blocks with the linear + nonlinear path design. The 2D convolution blocks cycle through dilation factors of 1, 2, 4, 8, 16, 32, 64 for four full passes with a total of 112 convolution layers per decoder. Decoders at lower levels receive input from the corresponding level of sequence representations, the interaction matrix prediction from one-level above and a 2D pairwise distance encoding matrix as an auxiliary input. 1D sequence representations are transformed to 2D with the pairwise sum operation ($Y_i = X_i + X_j$). The lower-level decoders predict for a subregion half the window size of the upper-level prediction, and the prediction from the upper-level corresponding to the window was upscaled by a factor of 2 and padded as input. For the distance encoding matrix $D$, for each cell type, $D_{ij}$ is the log distance-based expected balanced contact score at each genomic distance $|i−j|$ for intrachromosomal pairs ($i,j$), and interchromosomal pairs are filled with a constant of average interchromosomal log expected balanced contact score. The distance-based expectation scores for 32–256 Mb were monotonically transformed so that lower scores for lower distances are no higher than the shorter distances. The distance encoding matrix and the upscaled prediction from the upper level are combined with the 2D sequence representation by concatenation followed by a convolution block (Supplementary Fig. 1). The final model prediction is summarized by averaging with its transposes. The model predictions are averaged between the predictions from the forward- and reverse-complement sequences. The sequence encoder is also trained with an auxiliary task of predicting DNase-seq and ChIP-seq chromatin profile labels (Supplementary Table 6), which improved performance. To simultaneously predict chromatin profile labels and genome interactions, a 1D convolution block for predicting chromatin profiles is introduced which receives input from the 4-kb-resolution output of the sequence encoder.

Model training and evaluation. The processed micro-C datasets for H1-ESCs and HFF cells23 were downloaded from the 4D Nucleome (4DN) data portal (accession IDs 4DNF96GM12P8 and 4DNF634YP99). The genomic sequences were retrieved from the GRC38/hg38 reference genome. Training data were generated on-the-fly during training by uniformly sampling from training chromosomes with the Selene deep-learning sequence modeling library59. A separate model was trained for each micro-C dataset. The on-the-fly sampling generated new training samples for every training step. Each training sample consists of a sequence (the input) and the corresponding multilevel distance-normalized contact matrices (the target), which were also referred to as the genome interaction matrices. To compute the genome interaction matrices, the iterative correction matrix balance algorithm8 and adaptive propensity function were used to provide interaction matrices to the contact matrices retrieved from the micro-C datasets with cooler and cooltools packages60. Adaptive coarse graining is a preprocessing procedure implemented in the cooltools package that smooths the low-coverage areas of the contact map with adaptive window size and this step eliminates zeros by pooling the reads from the same position. No further smoothing was applied to the spatial resolution of the data. The processed matrix was then divided by the background matrix which is the exponential of the distance encoding matrix as described in the previous section (all operations are elementwise), and the minimum value of the background matrix was added to both nominator and denominator for numerical stability and noise reduction. The distance-based expectations are computed per chromosome with cooltools and then aggregated over all chromosomes.

The distance-expectation curve beyond a 1.6-Mb distance is smoothed with lowess. The chromosomes were divided into the training set (all chromosomes except for chr8, 9 and 10), the validation set (chr8) and the test set (chr9, 10).

The main loss function is the mean squared error between the predictions and the targets, or $\|\text{prediction} - \text{target}\|^2$, where prediction and target are both 250 × 250 square matrices, $N$ indicates the number of elements in the matrix to average over and the norm sign indicates the Frobenius norm. Missing values in the genome interaction matrices, which are typically due to low or no coverage, are ignored in the loss and gradient computation. An auxiliary binary cross-entropy loss function is also used to train the sequence encoding to simultaneously predict chromatin accessibility and ChIP-seq chromatin profile labels, or specifically the auxiliary loss is

$$\sum_{i,j} [\text{target}_{ij} \cdot \log(\text{prediction}_{ij}) + (1 - \text{target}_{ij}) \cdot \log(1 - \text{prediction}_{ij})]$$

where target$_{ij}$ is the binary chromatin profile target matrix of size $4 \times 250$ ($d$ is the number of chromatin profiles), and prediction$_{ij}$ is the predicted probability matrix of the same size, i and j are the indices of the matrices, and $N$ is the total number of elements in the matrix. The auxiliary loss is simultaneously trained on the same set of sequences as the main loss function. The list of chromatin profiles used is provided in Supplementary Table 6. The chromatin profile labels are generated for 4-kb bins and labeled one or zero based on whether any peak overlaps with the 4-kb bin.

To allow training of large-scale sequence models that do not fit into GPU memory with standard techniques, a horizontal checkpointing method was devised, leveraging the hierarchical structure of the model (Methods section ‘Scaling hierarchical deep-learning model training’ for details). Other training optimizations include parallelizing training data generation on CPU and randomly selecting either forward- or reverse-complement sequence for prediction, which can be seen as an unbiased stochastic approximation for averaging predictions from forward- and reverse-complement sequences.

For both flexibility in model application and efficiency in model training, the model was designed to be composed of three stackable modules (1Mb, 1–32 Mb, 32–256 Mb), which were trained in three stages. In the first stage, the sequence encoding at 4-kb resolution was pretrained with the task of predicting genome interactions within 1-Mb distance at 4-kb resolution and the auxiliary task of predicting chromatin profile labels at the same resolution (the cohesin-depleted HCT116 model was trained without auxiliary task). The encoder up to 4-kb resolution and the decoder trained in the first stage are also called Orca-1Mb. In the second stage, with the pretrained first section of the sequence encoder from the 1-Mb module, the multiscale 1–32-Mb model was trained to predict at 1-Mb, 2-Mb, 4-Mb, 8-Mb, 16-Mb and 32-Mb levels. For training multiscale prediction models, a series of subregions with increasingly smaller window size and finer resolution at each level, or the ‘zooming’ series, was selected. For example, for a 32-Mb sequence, a 16-Mb subregion was randomly selected, then an 8-Mb subregion within the 16-Mb region was randomly selected and this continued until a 1-Mb region was selected. The encoder up to 128-kb resolution and decoders trained in the second stage are also called Orca-32 Mb. In the third stage, the 32–256 Mb model is trained for both intrachromosomal and interchromosomal interactions, with the pretrained sequence encoder up to 128-kb resolution from the 1–32 Mb model. The full encoder and the third-stage decoders are also called Orca-32 Mb. The training data set for 32–256 Mb models were sampled from multiple chromosomes with the following process: a chromosome is first sampled, then add the full length of that chromosome to the sequence; then sample another chromosome, and add the full-length chromosome if not exceeding 256 Mb, and otherwise sample a subregion on that chromosome that makes up a total of 256 Mb. If the subregion continues adding other chromosomes until 256 Mb is reached, randomly permute the order of the sequence segments sampled and randomly select a strand direction for each segment; retrieve the corresponding sequence, intrachromosomal and interchromosomal genome interactions, and distance encodings as described above. The training process with stochastic gradient descent took about 150,000 steps for the first stage (1-Mb module was trained with learning rate 16, learning rate 0.002 with momentum 0.98 and the last 1/3 of steps are trained with stochastic weight averaging8), 150,000 steps for the second stage (32-Mb sequence with batch size 4 and learning rate 0.001 with momentum 0.98) and
20,000 steps for the third stage (256-Mb sequence with batch size 4 and learning rate 0.001 with momentum 0.98). The training hardware was one server equipped with four NVIDIA Tesla V100 32GB GPUs. The code for training Orca models with full details of the implementation is provided at the code repository.

Each training stage generates training data from micro-C data processed to different resolutions. The training data were sampled from the micro-C contact matrices at 1-kb resolution for the 1-Mb model, 4-kb resolution for the 1–32-Mb model and 32-kb for the 32–256-Mb model, and these high-resolution matrices are downsampled to the prediction resolution of the block outputs. Downsampling is performed by taking the average of the multiple entries that are collapsed into one, excluding the missing values. To further reduce overfitting, the input sequences for training are shifted by a random offset within 100 bp for the 1-Mb model, 1 kb for the 1–32-Mb model and 4 kb for the 32–256-Mb model.

Model prediction evaluation on holdout test chromosomes. To evaluate the model prediction performance on holdout test chromosomes, multiscale genome interaction matrices were systematically predicted on the test chromosomes and the predictions were compared with the observed micro-C data. The evaluation data were processed in the same procedure as for training data generation. Missing values in the micro-C target matrices are excluded from the evaluation (missing values are typically due to low or no coverage). Because target matrices at lower resolutions are downsampled from higher-resolution matrices by the binning procedure described above, a downsampled value is computed from averaging multiple values from high-resolution matrix while excluding missing values, and if 25% of these values are missing then the targeted value is excluded in evaluation. Specifically, for evaluating the predictions at 1–32-Mb levels, the test set test chromosomes were tiled with 32-Mb windows at a step size of 0.5 Mb. For each 32-Mb window the genome interactions were predicted at all scales from 1 Mb to 32 Mb by sequentially zooming into 16-Mb, 8-Mb, 4-Mb, 2-Mb, 1-Mb subwindows each located at the center of the higher-level region. All prediction matrices were then completed and flattened, multiple random disruptions were introduced between the predictions and micro-C observations. The 1-Mb level performance of the 1–32-Mb models was also compared with the 1-Mb module predictions on the same 1-Mb windows.

For evaluating the intrachromosomal 32–256-Mb-scale predictions, two 256-Mb sequences each containing a test chromosome were first generated with the rest of the 256-Mb length padded with sequence from chr1 (only the intrachromosomal interactions were evaluated). For predictions at 128-Mb, 64-Mb and 32-Mb levels, the same starting positions that tile the test chromosomes with step size of 5,120 kb were used. Windows that extended beyond the test chromosome were excluded because of missing values in the micro-C data. To evaluate the predictions at 10-bp level were used. The Orca 1-Mb-level predictions and target genome interaction matrices were re-used by linearly upsampling with a factor of 2 and cropped to the Akita output region, and additional Gaussian filtering with sigma 1 and kernel size 5 and clipping to (−2, 2) was then applied to match the Akita data-processing steps. For each test sample, background-subtracted Pearson correlations were computed against the Akita targets and Orca targets processed as described above. To compute the background-subtracted Pearson correlation for a target matrix, each score was subtracted by the average scores at the same distance in the same matrix before computing correlation. The background subtraction has minimal effects on preserving the genome structure information and improves robustness to different data preprocessing.

Scaling hierarchical deep-learning model training. To scale deep-learning sequence models to hundreds of megabases, a scalable memory-efficient training algorithm was devised to dramatically reduce the memory requirement. As illustrated in Supplementary Fig. 25, the regular training procedure for deep learning is layer-wise and stores all internal representations in memory for computing gradients and thereby results in extreme memory demand for large model input. Checkpointing is a memory-saving technique first developed for residual networks with a high number of layers. With checkpointing, only internal representations at the checkpoint layers are stored and other internal representations can be recomputed on-the-fly when gradient computation is needed. However, even with the checkpointing technique, training is still infeasible for very large sequence input because the memory requirement of computing even only the first layer for a single sequence is beyond the maximum capacity of currently available GPUs.

By leveraging the hierarchical structure of the sequence model, the memory consumption of the bottom layers, which use the most memory, can be greatly reduced by executing them in horizontal blocks and only storing the output of the blocks. This approach fixed the memory usage of the lower layers to the memory needed to compute the block, with the minimum block size being the receptive field of the block output layer (recommended sizes are at least double of the minimum for computational efficiency). For example, the receptive field of the 4-kb-resolution layer output of the Orca sequence encoder is 212 kb, which is less than 1/150 of 32 Mb or 1/1,200 of 256 Mb, allowing great reduction of memory usage. Because the memory consumption in the bottom layers is orders of magnitude larger compared with the upper layers, this essentially resolved the memory consumption issue. The Orca models and all those multiresolution models beyond whole-chromosome-scale input. I refer to this technique as horizontal checkpointing. Horizontal checkpointing was used to allow the model to scale to large input for training and prediction of Orca 32-Mb and Orca 256-Mb models. Horizontal checkpointing also allows gradient computation during model training, and while this capability was not utilized in the current models due to the increased training time, such capability could be useful in future studies.

SV impact on multiscale genome interactions. Orca models allow the prediction of the multiscale genome organization impact of almost any genome variant at any predefined resolution of the whole genome sequence. This is a unique value since the available sample value was not previously available.

To evaluate the prediction performance of Orca models, we constructed a set of synthetic variants in a multiresolution fashion to represent the structural impact of their disruption. To perform a genome-scale screening, the analysis was sped up by introducing a multiplexed approach to in silico mutagenesis. Since 10-bp sequences with strong structural impacts are sparse (most disruptions have near zero effects), multiple random disruptions can be introduced to the same sequence with a very low probability that more than one disruption will have a strong effect. The multiplexed design ensures that for the effects of a large number of mutations that cover the genome was designed. In this study, a score was assigned to all genomic sequences in 10-bp bins on autosomes representing the structural impact of their disruption. To perform a genome-scale screening, the analysis was sped up by introducing a multiplexed approach to in silico mutagenesis. Since 10-bp sequences with strong structural impacts are sparse (most disruptions have near zero effects), multiple random disruptions can be introduced to the same sequence with a very low probability that more than one disruption will have a strong effect. The multiplexed design ensures that for each 10-bp sequence, multiple random disruptions are introduced in different sequences, each with a different set of random disruptions. The 10-bp site-specific sequence disruption effect is measured by the absolute score of all sequences that carry a disruption of the 10-bp sequence. The disruption impact on local genome interactions is measured by 1-Mb structural impact score, which is the average absolute log fold change of interactions between the disruption position and all other positions in the 1-Mb window. The Orca 1-Mb modules for H1-ESCs and HFFs are used for all predictions to allow fast screening of a large number of sequences.

More specifically, the genome was tiled with 1-Mb windows at 0.8-Mb step size across all autosomes. Each 1-Mb window is considered as a 25 × 40-kb region containing 4,000 × 10-disruption sites. Thus, 12,000 mutated sequences are generated for each 1-Mb window. Each generated sequence contains 20 disruptions in each 25 × 40-kb region. Each 10-bp sequence disruption matrices was multiplexed to 10-bp sequences to a 20 × 4,000 matrix with each row containing all 10-bp sites of a 40-kb region, then randomly shuffling each row independently, resulting in 4,000 columns each corresponding to a mutated sequence. This process was repeated three times to generate 12,000 sequence designs. According to these designs, 10-bp sequence disruption matrices were generated by linearly upsampling the Orca sequence encoder predictions with a factor of 2 and cropping to the Akita output region, and additional Gaussian filtering with sigma 1 and kernel size 5 and clipping to (−2, 2) was then applied to match the Akita data-processing steps. To compute the background-subtracted Pearson correlation for a target matrix, each score was subtracted by the average scores at the same distance in the same matrix before computing correlation. The background subtraction has minimal effects on preserving the genome structure information and improves robustness to different data preprocessing.
sequence disruptions are introduced by replacing the original 10-bp sequence with random nucleotides that match the nucleotide composition in the 1-Mb window. For motif enrichment analysis, vertebrate nonredundant motifs were downloaded from the ZFIN database. Motif matches for each 10-bp flanking search were scanned for extending by 10-bp flanking sequence on each side, and a maximum log-odds score over the 30-bp window for each motif was obtained. The maximum motif log-odds score in this window was also referred to as the maximum motif log-odds score for the 10-bp site. To avoid overlap of extended sequences, only one 10-bp site every 30-bp was considered for statistical tests. To analyze non-CTCF motif enrichments, 10-bp sites with 1-Mb structural impact score > 0.01 and without nearby CTCF motif matches (CTCF max motif log-odds < 6 within 200bp) or CTCF binding sites (CTCF ChIP-seq fold over control < 4; ENCODE accession IDs ENCF4731ZV, ENCF7619HS) were used. Next, to quantify the enrichment of motifs, two-sided t-test (without assuming equal variance) was performed to compare the motif log-odds scores of these filtered sites against the background of 100,000 randomly drawn motifs. Fold enrichment was also computed on the same motif with a motif log-odds threshold of 12.

For motif enrichment analysis of the chromatin compartment activities, two virtual screens were designed. For each screen, a pool of source sequences and one or more target positions were selected. For every source sequence and target location pair, the source sequence was ‘inserted’ into the target location by swapping out the original sequence at the target position, then the genome interaction pattern changes were predicted and quantified by the 32-Mb structural impact score (the average absolute log fold change of interactions between the target position and all other positions in the 32-Mb window). Because a large proportion of the mutated sequence is in common with the original sequence, the computation was sped up by only recomputing the internal representations that are affected by the change.

The source sequences were generated from a large genomic region or across entire chromosomes by dividing the region into fixed-sized segments, and the structural impact scores of the source sequences at all positions were visualized as a chromatin compartment alteration activity profile. For exploratory virtual screens, a 32-Mb region, chr10:77,072,000–109,072,000 covering multiple A compartment, B compartment and intermediate regions, was used. Activities of source sequences tiling this region were screened for at nine target positions from the large-scale screen of 200 targets, by taking the first principal component across the 200 compartment activity profiles (one for each target position). The sign of a principal component is arbitrary, but the direction that corresponds to compartment A activity can be easily detected, such as based on TSS enrichment. The top 2% of sequences with the strongest compartment A activity were used for downstream enrichment analysis.

For enrichment analysis of the chromatin compartment activities, the FANTOM CAGE signal profile (maximum count across samples) was downloaded from the UCSC table browser with a filter of count > 1, and the annotations for TSS, 5' UTR, 3' UTR, exon and genes were from Ensembl release 97. The chromatin state annotations for HCT116 are from EpiMap.

For performing analyses with random permutation of sequences, the sequence to be permuted was first divided into segments of the same specified length, then the order of the sequence segments was randomly permuted. As random permutation disrupts any sequence patterns larger than the segment length, this analysis can be used to reveal the length scale of the sequence dependencies.

**Reporting Summary**
Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

The GRCh38/hg38 reference genome and 3D genome datasets under 4DN accession numbers 4DNF69GMP218, 4DNF66450Y1P and 4DNFILP999JS were used for training the Orca sequence models. All coordinates in the manuscript refer to GRCh38/hg38 unless otherwise indicated. SV experimental validation datasets were downloaded from NCBI GEO accesses GSE137372, GSE562835, GSE78109 and EBI ENA accession PRJEB5236. Data used and generated in this manuscript were also deposited into Zenodo: https://zenodo.org/record/6234936, https://zenodo.org/record/4594676 and https://zenodo.org/record/6227750.

**Code availability**

All code, models and data for running Orca are available from the Github repository https://github.com/zhoulab/orca (https://doi.org/10.5281/zenodo.6257290). A user-friendly web server is available at https://orca.zhoulab.io. The manuscript analysis code is available at https://github.com/zhoulab/orca_manuscript (https://doi.org/10.5281/zenodo.6257292).

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**Author contributions**

J.Z. conceived and designed the study, developed the computational methods, performed the analysis and wrote the manuscript.

**Competing interests**

The author declares no competing interests.

**Additional information**

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Extended Data Fig. 1 | Performance of Orca model predictions for the HFF cell type. a). A multiscale sequence-based prediction example zooming from whole-chromosome into a position on a holdout test chromosome. Predictions from 1–256 Mb scales are compared with micro-C experimental observations. Missing values in micro-C data are shown in gray, and these regions are also indicated in the 64–256 Mb prediction heatmaps because predictions at major assembly gaps or unmappable regions are of unknown accuracy. The genome interactions are represented by the log fold over genomic-distance-based background scores for both prediction and experimental data. b). Scatter plot comparison of the predicted interaction scores with the micro-C measured interaction scores (log fold over background) on the holdout test chromosomes. 10,000 randomly subsampled scores are shown in each panel. The overall Pearson correlations across the entire test chromosomes are annotated. The genome interactions are represented by the log fold over background scores for both prediction and experimental data. Predictions for 1–32 Mb levels are from the Orca-32Mb model and 64–256 Mb levels are from the Orca-256Mb model.
Extended Data Fig. 2 | Performance of Orca model predictions for cross-cell-type genome interaction difference. a). Scatter plot comparison of the predicted cell type differences of genome interactions (H1-ESC - HFF) with the micro-C measured interaction score differences on the holdout chromosomes. 10,000 randomly subsampled scores are shown in each panel. The overall Pearson correlations across the entire test chromosomes are annotated. The genome interactions are represented by the log fold over genomic-distance-based background scores for both prediction and experimental data. b). Prediction performance for position pairs with the strongest absolute log-fold differences between the two cell types (top 1 percentile). The performance of models predicting the cell type labels (the cell type with stronger interaction) is measured by receiver operating characteristic (ROC) curve. The area under the ROC curve (AUROC) is annotated. The AUROC score can be interpreted as the probability of a randomly selected positive example (that is stronger in HFF) being ranked higher than a randomly selected example (that is stronger in H1-ESC). Predictions for 1–32 Mb levels are from the Orca-32Mb models and 64–256 Mb levels are from the Orca-256Mb models.
Extended Data Fig. 3 | Example Orca predictions of Polycomb-mediated interactions. Predicted and observed H1-ESC and HFF genome interactions for two regions from a holdout chromosome, a) chr10:116850000-117850000 and b) chr10:100450000-101450000 are shown. The predicted and observed Polycomb-mediated interactions are marked with black triangles. ChIP-seq signal tracks for CTCF and H3K27me3 for the two cell types are also shown. Polycomb-mediated interactions are predicted to be specific to H1-ESC in both examples, consistent with experimental micro-C and ChIP-seq data.
Extended Data Fig. 4 | Example Orca predictions of promoter-enhancer interactions. Predicted and observed H1-ESC and HFF genome interactions for two regions from holdout chromosomes, a) chr8:127400000-128400000 and b) chr9:94360000-95360000 are shown. The predicted and observed enhancer-promoter interactions are marked (promoter positions or promoter-promoter interactions are marked with red triangles, enhancer-promoter or enhancer-enhancer interactions are marked with black triangles; we only marked a subset of all interactions observed). ChIP-seq signal tracks for CTCF and H3K4me3, H3K27ac, and H3K4me1 for the two cell types are also shown. The predicted enhancer-promoter interactions are consistent with micro-C observations and enhancer histone mark signal from ChIP-seq data.
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | Visualized predictions of transposon-mediated boundary element insertion effects in multiple insertion sites. All insertions with previously categorized effects (boundary creation, boundary strengthening, and no domain-level effect) in Zhang et al.24 are shown. The experimental measurements by in situ Hi-C in HAP1 cell is compared with Hi-ESC model predictions. The genome interactions are represented by the log fold over genomic-distance-based background scores for both prediction and experimental data. Arrows indicate the insertion sites. The genome coordinates are in hg19.
Extended Data Fig. 6 | Comparison of Orca prediction with Capture Hi-C experimental measurement for structural variants from Franke et al. 2016.
Capture Hi-C data from mouse with SVs are compared with predictions for effects of equivalent human structural variants. Predicted log fold over background at 4 Mb level are scaled with the distance-expectation curve from capture Hi-C.
**Extended Data Fig. 7 | Multiplexed in silico mutagenesis screen results are highly correlated with single-mutation in silico mutagenesis screen results.**

**a.** Predicted structural impact scores (1 Mb) of single disruptions (left) and multiplexed disruptions are shown on the y-axis, with disruption positions on the x-axis. 10 bp disruption sites screened cover the center 0.8 Mb of the 1 Mb region. The first three rows are three independent runs (for single disruption only the disrupted sequences are random across the runs, and for multiplexed disruption both the multiplex design of disruption sites and the disrupted sequences are random), and the last row shows the minimum of the three at each position. **b.** Relationship between the correlation of single and multiplexed disruption profiles (y-axis) and the number of runs combined (x-axis).
Extended Data Fig. 8 | Visualization of virtual screen sequence activity on chromatin compartment alteration. A subset of 1000 contiguous source sequences among all 27981 12800 bp source sequences covering chr8, 9, and 10 are shown. Target locations are ordered by the main mode of compartment change detected at the target site (from top: A>B to bottom: B>A), which is quantified by the loading of the first principal component of the whole sequence structural impact score (32 Mb) matrix.
Extended Data Fig. 9 | Random sequence permutation effects on sequence compartment A and compartment B activity. Comparison of chromatin compartment activities of 25600 bp sequences permuted by different segment length (at each permutation segment length, 2bp, 4bp, …, 256bp, every 25600 bp sequence is divided into segments and the segments are then randomly shuffled and concatenated). Compartment B activity is compared with sequence A/T content at the same locations.
Extended Data Fig. 10 | Predicted effects of disrupting genomic regions by randomly permuting sequences. At each disruption site indicated by the arrow, 1.28 Mb sequence centered at the position is permuted by 4 bp segments. Permutated compartment A sequences show B compartment interaction patterns, while disrupted compartment B sequences remain to be in B compartment.
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- Data collection: No software was used
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The GRCh38/hg38 reference genome and 3D genome datasets under 4DN accession numbers 4DNIF9GMP2J8, 4DNIF6430YPP9, 4DNIFLP99Q5S were used for training the Orca sequence models. All coordinates in the manuscript refer to GRCh38/hg38 unless otherwise indicated. SVs experimental validation datasets were downloaded from NCBI GEO accessions GSE137372, GSE66383, GSE78109 and EBI ENA accession PRJEB5236. Data used and generated in this manuscript were also deposited into Zenodo repositories https://zenodo.org/record/6234936, https://zenodo.org/record/4594676, and https://zenodo.org/record/6245470.
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Sample size  No sample size calculation was performed. We used the highest sample size possible whenever feasible e.g. all 10bp sites in the genome and such sample was sufficient for generating the statistically significant results reported in the manuscript.

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