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4D genome rewiring during oncogene induced and replicative senescence

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SUMMARY

While transcriptional and epigenetic changes associated with senescence are well studied, the role of the extensive senescence-associated 3D genome reorganization remains elusive. Here, we have generated genome wide chromatin interaction maps, epigenetic, replication-timing, whole genome bisulfite sequencing and gene expression profiles from cells entering replicative senescence (RS) or upon oncogene induced senescence (OIS). We identify Senescence Associated Heterochromatin Domains (SAHDs). Differential intra vs inter SAHD interactions lead to the formation of senescence associated heterochromatin foci (SAHFs) in OIS but not in RS. This OIS-specific configuration brings active genes located in genomic regions adjacent to SAHDs in close spatial proximity and favours their expression. Finally, screening of factors for SAHF induction revealed DNMT1 as a novel component that induces SAHFs by promoting HMGA2 expression. Upon DNMT1 depletion, OIS cells transition to a 3D genome conformation akin to that of cells in replicative senescence. These data show how multi-omics and imaging can identify critical features of RS and OIS and discover new determinants of acute senescence and SAHF formation.

Keywords: Senescence, replicative senescence, oncogene induced senescence, 3D genome architecture, Hi-C, chromatin compartments, gene regulation, DNMT1

Highlights

- Deep multi-omics characterization of replicative and oncogene-induced senescence
- Senescence-associated heterochromatin domains (SAHDs) form SAHFs via 3D changes
- DNMT1 is required for SAHF formation via regulation of HMGA2 expression
- SAHF formation leads to expression of SAHF-adjacent genes via 3D chromatin contacts

INTRODUCTION

Senescence was first described in normal human fibroblasts as a permanent proliferation arrest due to the exhaustion of the cellular replicative potential, termed replicative senescence (RS) (Hayflick, 1965; Hayflick and Moorhead, 1961). Subsequently, it was demonstrated to occur in primary cells in response to oncogene activation and, in that case, it was named oncogene induced senescence (OIS) (Serrano et al., 1997). Over the past two decades, the concept of senescence was revisited and further extended to incorporate processes leading to cell cycle arrest in response to a variety of insults, including DNA damage, oxidative stress and chemotherapeutic drugs (Campisi and d'Adda di Fagagna, 2007; Di Micco et al., 2011; Kuilman et al., 2008; Narita et al., 2003; Schmitt et al., 2002), as well as to specific stress signals in embryo development during morphogenesis and in wound healing and regeneration (Munoz-Espin et al., 2013; Storer et al., 2013; Yun et al., 2015). All these events are commonly referred to as premature senescence. Therefore, senescence is now considered to be a stress response program.
RS and OIS cells display a variety of common features, like accumulation of senescence-associated beta-galactosidase (SA-β-gal), activation of Rb/p16 and p53/p21 pathways, morphological changes and induction of the senescence-associated secretory phenotype (SASP) (Campisi, 2013). However, they differ markedly in their nuclear architecture. On the one hand, OIS nuclei are highly compartmentalised. They display heterochromatin bodies, called senescence-associated heterochromatin foci (SAHF), enriched in H3K9me3 and other core heterochromatin marks in different human cell types and pathologic conditions (Chandra et al., 2012; Narita et al., 2003; Sun et al., 2018; Xu et al., 2014). RS nuclei on the other hand are enlarged and display a variety of features, ranging from compaction of individual chromosome arms to distension of peri-centromeric regions (Cruickshanks et al., 2013; De Cecco et al., 2013; Swanson et al., 2013).

3D genome organisation provides a link between various cellular processes such as DNA replication, transcription, DNA repair and gene expression (Bonev and Cavalli, 2016). Recent studies employing microscopy (Cremer et al., 2015) and chromosome conformation capture based methods like Hi-C (Sati and Cavalli, 2017) have improved our understanding of genome organisation. Hi-C methods have revealed a hierarchy in genome organisation, where the basic units of genome folding, represented by TADs (Topologically Associating Domains), are organised into two compartments: the early replicating or active (A) compartment and the late replicating or inactive (B) compartment. These compartments then coalesce into chromosome territories (Bonev et al., 2017; Dixon et al., 2012; Nora et al., 2012; Sexton et al., 2012). Likewise, microscopy approaches have identified chromatin domains (CDs) which might represent the microscopical counterpart of TADs. CDs form clusters that can belong to either the inactive or the active nuclear compartment and together they form chromosome territories. Microscopy also identified a so-called interchromatin compartment (IC), made of channels that separate chromatin domains and is particularly relevant for gene regulation since active genes and soluble chromatin regulatory components tend to locate at the surface and within the IC (Cremer et al., 2015). Furthermore, single nucleosome imaging methods support this model as they found transcription centres in the IC regions and transcriptional hubs restricting chromatin mobility (Nagashima et al., 2019).

While all these approaches have advanced our understanding of 3D genome and nuclear organization, the relation between microscopical observations and chromatin contacts remains largely unexplored and the functional relevance of observations made with both approaches needs to be studied. In particular, our knowledge about higher-order chromatin changes during different types of senescence is limited. Very few studies employing Hi-C assays have been undertaken to compare 3D changes associated with RS or OIS nuclei (Chandra et al., 2015; Criscione et al., 2016). One such study, employing low resolution Hi-C on early stages of OIS identified no changes in TAD borders (Chandra et al., 2015), whereas another study performed on RS cells displayed changes in TAD borders and compartmental switching of some TADs from active to inactive compartments and vice versa (Criscione et al., 2016; Zirkel et al., 2018). These studies found a shift in the ratio between short- and long-range chromatin contacts, but they were not in agreement on the direction of this shift. In addition, it is still unknown whether there are differences between various chromatin architectural properties like insulation, compartmentalisation and transcription start site interactions in RS or OIS nuclei, or if there is any intermediate chromatin folding stage that might be common between RS and OIS. The relationship between the 3D genome architecture, its underlying epigenome and gene expression has never been compared between RS and OIS, and differential interactions among the H3K9me3 marked heterochromatin domains, that form SAHF in OIS but not in RS, might be of major importance in these regulations.
In this study, we have addressed these issues by utilizing human fibroblast-based RS and OIS systems to understand similarities and differences in their 3D genome organisation. We performed Hi-C, ChIP-Seq for key heterochromatin / euchromatin marks, replication timing experiments, whole-genome bisulfite sequencing and RNA Sequencing in OIS and RS. This enabled us to delineate an interplay between 3D architecture and the underlying epigenetic landscape on the one hand, and transcriptional changes on the other. Our observations with omics approaches were validated using oligopaint-based 3D DNA FISH assay. Based on the observations learned from this analysis, we tested various causative scenarios. Finally, we have identified DNMT1 as a new factor that causes senescent 3D genome reprogramming and its difference between RS and OIS.

RESULTS
Differences in 3D genome organisation in OIS and RS are based on distinct interactions among chromatin compartments.

WI38 primary fibroblasts (RS-Proliferative) and their serial passaging-induced replicative senescent stage (RS-Senescent) were used as a RS system (Figure 1A). For OIS, we used the WI-38hTERT/GFP-RAF1-ER (referred to as “RAF cells”) model system (Jeanblanc et al., 2012), where OIS was induced by adding 4-hydroxytamoxifen into the culture media. Following RAF induction, the cells progress into senescence, triggering cell cycle arrest by day 2 (OIS-D2), and form SAHFs in almost all cells by day 4 (OIS-D4) (Figure 1A). In contrast, a few bright DAPI foci appear in RS, but no SAHFs are observed (Figure 1A). To follow the appearance of the cell senescence phenotype, we performed a classical SA-βGal assay in OIS and RS. From OIS-D4 onwards all cells display SAHFs. By OIS-D6 all cells are senescent, and they remain in senescence when followed up to OIS-D10 (Figure S1A-B). We further confirmed the senescence status by bromodeoxyuridine (BrdU) incorporation in both RS and OIS (Figure S1C) and, as expected, both RS-Proliferative and RAF cells were found to be karyotypically normal (Figure S1D). To get a homogenous population for Hi-C, cells were FACS sorted, selecting for the G1 phase (Figure S1E) and Hi-C was performed in replicates on non-induced cells (OIS-D0), RAF-induced cells after 2, 4, 6 and 10 days of induction (OIS-D2 to OIS-D10), RS-Proliferative and RS-Senescent cells. In total, 4.5 billion Hi-C contacts were generated and analysed in this study (Table S1).

The normalised Hi-C matrices displayed a progressive global reorganisation in both OIS and RS nuclei. OIS displayed a huge gain in long-range contacts, which became prominent by OIS-D4 (by which time SAHFs are formed) (Figure 1B). Hi-C matrices of OIS-D6 and OIS-D10 are visually similar regarding long range contacts, suggesting that a steady-state in chromatin architecture is reached starting from OIS-D6 onwards. Strikingly, the mean intrachromosomal contact probability over distance displays a gradual shift from close-cis (<260kb) to far-cis (>30Mb) in both OIS and RS (Figure 1C). For intermediate distances (260 kb – 2Mb), corresponding to most of the large chromatin domains or TADs, RS nuclei display more cis contacts that their proliferative counterparts (Figure 1C, Figure S1F). This gain in long-range contact suggests chromosome compaction, which was confirmed in DNA-FISH experiments by a decrease in whole-chromosome area in OIS-D6 versus OIS-D0, as well as RS-Senescent versus RS-Proliferative nuclei (Figure 1D-E). To compare this increase in long range contacts other cell type we also generated Hi-C maps from BJ-hTERT-B-RAF-V600E (henceforth called BJ-raf) and from skin fibroblasts form a 74 year- old human (HSF74). Since HSF74 cells were still growing, they were passaged into replicative senescence and Hi-C maps were produced from both HSF74-Proliferative and HSF74-Senescent cells (Figure S1G-K). We further compared our chromosomal contact probability profiles with previously published
OIS and RS datasets (Figure S1K-L). Except for HUVEC cells, changes in long range contacts observed in our OIS and RS system were in concordance with previously studied cell types (Figure S1L) (Chandra et al., 2015; Zirkel et al., 2018).

Since the previous studies were performed with different cell types, for downstream comparisons we focused on the WI38-based OIS and RS systems, which share the same cell background. Analysis of Hi-C profiles shows that, in OIS, the regions gaining long-range interactions are in the B compartment (Figure 1B, Figure S1M-N). To quantify the changes in compartment interactions, we identified TADs and classified them into two compartments: active (A) and inactive (B) (see STAR methods). Plotting the log ratio of observed versus expected contacts revealed a progressive increase in compartmentalisation during OIS, with a preferential gain of B-B interactions and loss of A-B interactions (Figure 1F). In contrast, RS leads to a loss of A-A interactions and a preferential gain of A-B interactions that highlights a decrease in genome compartmentalisation (Figure 1F). We confirmed these results by using an alternative approach based on eigenvector values (see STAR methods) (Figure 1G).

In conclusion, senescence is associated with a major 3D genome reorganisation, with progressive loss of close-cis and gain of far-cis contacts. However, RS and OIS differ markedly in their organisation of genomic compartments. RS cells reduce compartmentalisation due to a relative decrease in A compartment interactions, whereas in OIS, the genome organisation is dominated by heterochromatin or B-B interactions leading to strong genome compartmentalisation, which is evident in the form of SAHFs.

**Long-range interactions among Senescence Associated Heterochromatin Domains (SAHD) generate SAHFs**

To identify the regions involved in SAHF formation, we combined epigenome profiling with diffHic analysis (Lun and Smyth, 2015). The systematic overlay of differentially interacting regions from diffHic with 27 different histone modifications from IMR90 cells (NIH roadmap to epigenomics) in the WashU browser (Lun and Smyth, 2015) highlighted broad H3K9me3 enriched regions showing a relative gain in long-range interactions in both RS and OIS (Figure 2A) (Zhou et al., 2013; Zhou et al., 2011). We thus performed ChIP-Seq experiments for H3K9me3, H3K27me3 and H3K4me3 on OIS and RS cells. Data analysis showed these identified regions to be selectively enriched for H3K9me3, but not for other marks, both in OIS and in RS (Figure S2A-B). We defined these conserved OIS H3K9me3 regions as Senescence Associated Heterochromatin Domains (SAHDs) (Table S2). We found SAHDs to be gene poor, enriched in constitutive LADs, L1 and L2 isochores, LINE / LTR elements, poor in SINE elements / simple repeats and, interestingly, enriched in late replication timing regions (S4 and G2), (Figure S2C-G).

The quantification of inter SAHD interactions showed that OIS cells display a much larger gain compared to RS cells. Furthermore, OIS cells massively lose intra SAHD interactions, while the effect is moderate in RS (Figure 2B and S2H). Interestingly, SAHDs also gain inter SAHD interactions in trans in OIS conditions (Figure S2I). Thus, SAHDs are a general feature of senescent cells and the formation of SAHFs only in OIS cells correlates with a shift from intra- to inter-SAHF interactions that is more prominent than in RS. To test this hypothesis, we performed 3D-FISH in a selected SAHD subset. Since the regions of interest (SAHDs) were large in terms of genomic size (1 Mb – 4 Mb), we employed an oligopaint-based approach for the FISH assay (Figure 2C-E, Figure S2J, Table S3). Image analysis of OIS-D0 and OIS-D6 FISH data showed that SAHDs (probes A, B and C) localize within SAHFs upon OIS (STAR Methods) (Figure 2D, F). The appearance of SAHFs in OIS-D6 occurred concurrently with
the relocation of SAHD probes away from the nuclear periphery, resulting in strong shortening of the inter-SAHD 3D distances (Figure 2G-H). Size normalised inter SAHD distances were constant with these observations (Figure S2K). Furthermore, SAHDs display an increase in the signal diameter in OIS, consistent with internal chromatin decompaction and reduction of intra-SAHD contacts (Figure 2I). In contrast, image analysis of RS-Proliferative and RS-Senescent FISH data indicates partial displacement of SAHD probes from the nuclear periphery and no significant change in the inter SAHD distances (Figure 2E, 2J-K). However, size normalisation in RS-Senescent condition display a significant decrease in all inter probe distances, which is in concordance with our previous observation of massive chromosome compaction in RS-Senescent cells (Figure 1E, S2K). Moreover, not all SAHD probes display an increase in area in RS cells and only SAHD probes that move away from the periphery (B and C) display an increase in signal diameter (Figure 2J, L), suggesting that movement from the nuclear periphery towards the interior might be associated with SAHD decompaction. Finally, we found SAHFs to be conserved in BJ cells and, like in WI38 OIS cells, SAHDs in BJ cells also loose intra-SAHD and gain inter-SAHD interactions under OIS conditions (Figure S2L-M). FISH on one of the SAHDs (Probe A) confirm localization into SAHFs in raf induced OIS in BJ cells (Figure 2M-N). Furthermore, SAHDs were displaced from the nuclear periphery and displayed an increase in signal diameter in OIS (Figure 2O-Q).

Hi-C matrices display part of the 3D chromosome information in 2D matrices. Previous work has shown that these matrices can be analysed by the TADbit modelling tool to derive 3D chromosome folding models that are in good agreement with in vivo chromosome architecture (STAR Methods) (Bau et al., 2011; Mas et al., 2018; Serra et al., 2017). We thus used TADbit to model all chromosomes from OIS-D0 and OIS-D6 conditions at 100kb resolution. We then quantified the distance distribution between SAHD and non-SAHD regions on chromosome 5 and compared them with 3D FISH data. This analysis shows that the distance distribution from our 3D models is in good agreement with experimental 3D FISH data and provide means of visually inspect the interaction data in 3D (Figure S2N-O). In conclusion, deep Hi-C sequencing leads to the genome-wide identification of the chromosomal domains, called SAHDs, that form SAHFs upon oncogene induction. These regions display unique architecture in terms of epigenetic marks, replication timing, sequence composition and 3D organization. Their 3D models along with the Hi-C maps and relative ChIP-seq tracks can be visualised in the MuGVRE browser. (https://vre.multiscalegenomics.eu/data_repositories/data_senescence.php).

Detachment from the nuclear lamina and weakening of inter SAHD interactions can lead to SAHF formation

Previous studies suggested that SAHF formation occurs in OIS due to lamina degradation, causing release of heterochromatin from the lamina which, in turn, was hypothesized to lead to heterochromatin aggregation (Sadaie et al., 2013; Shah et al., 2013). Detachment of constitutive lamina-associated domains (cLADs) in OIS has been recently confirmed but, surprisingly, knockout of the Lamin B receptor (LBR) does not induce detachment of cLADs (Lenain et al., 2017). These data suggest that other forces can maintain heterochromatin localization at the nuclear periphery in the absence of the LBR anchor. Intriguingly, Hi-C and FISH analysis showed that the movement of SAHDs away from periphery was correlated with SAHD decompaction. Therefore, we decided to quantitatively assess the relative roles of SAHD chromatin decompaction and release of from the nuclear periphery in SAHF formation. To this aim, we built a polymer model of an 80 Mbp region of the q-arm of chromosome 5, which encompasses FISH probes B, C and D used in this study. Chromatin was modelled as
self-avoiding block copolymer moving in a cubic box (STAR Methods) (Figure 3A). We considered that SAHDs may self-attract at short-range with strength $\varepsilon$ and that cLAD regions may transiently interact with the nuclear membrane with strength $\gamma$. We first inferred model parameters ($\varepsilon = -0.055kT$, $\gamma = -0.6kT$) to quantitatively describe SAHD organization in cycling/D0 cells (STAR Methods) with the formation of medium-size droplets spread in the simulation box, typical of micro-phase separation (Ghosh and Jost, 2018; Jost et al., 2014) (Figure 3B-D, S3A-C). This model was then used to test the effects of different hypotheses for how chromatin alterations could induce the observed 3D genome organization. Initially, we tested the two following cases: 1) a time relaxation (TR) scenario, in which we simulated 6 days of real time without modifying model parameters; 2) a membrane release (MR) scenario: after 24 hours of real time, we imposed a detachment of SAHDs from the membrane periphery (Figure 3E-G, S3D-F). These predictions are not consistent with the experimental data where SAHDs locate internally and SAHD contacts at OIS/D6 show a weak decrease ($\sim 10\%$) at small scales ($< 1 - 5$ Mbp) and a stronger increase ($\sim 20-30\%$) at larger scales ($>10-20$Mbp), compared to OIS/D0. In the MR scenario, we observed a very weak decrease in relative contacts ($\sim 1\%$) at small scales and a strong increase at intermediate and large scales ($\sim 35-50\%$) at OIS/D6 compared to OIS/D0 (Figure 3M-J, S3J-I). SAHDs now form large, SAHF-like, compartments localized away from the membrane (Figure 3H-J, S3G-I). The formation of SAHF in the MR scenario is more compatible with experimental observations than the TR scenario. However, the predicted rise in relative SAHD contacts is too strong, compared with the experimental Hi-C data. Since both Hi-C and DNA FISH show a decondensation of SAHDs during senescence (Figure 2B, 2I), we tested the combination of the MR scenario with a weakening of the interaction strength (reducing the value of $\varepsilon$) between heterochromatin monomers (called MR-WI scenario). We found that lowering $\varepsilon$ to -0.035 kT leads to consistent predictions at all length and time scales, with a detachment of SAHDs from the membrane and formation of SAHF (Figure 3K-M, S3J-L). Furthermore, a comparison of the MR-WI scenario with DNA FISH data showed that the model can reproduce quantitatively the changes in distance between SAHD (B, C) and non-SAHD (D) probes (Figure 3N). Finally, we compared the top-down, constraint-based chromosome model (produced by TADbit) with the bottom-up approach using the MR-WI scenario. The two models show a positive correlation, both in global chromosome conformation and in specific 3D folding of SAHD regions (Figure S3N-O).

Taken together, these data suggest that local decompaction of SAHDs and their detachment from the nuclear lamina can explain quantitatively SAHF formation observed in OIS senescence and the genome-wide changes in Hi-C contacts.

**Identification of DNMT1 as a new component required for SAHF formation**

Since SAHDs correspond to genomic regions that replicate late in proliferating cells, we tested whether replication timing is perturbed upon OIS. For this, we performed replication timing sequencing (Repli-Seq) on un-induced RAF cells (OIS-D0) and RAF cells after 12 hr, 24hr and 36 hr of RAF induction. The Repli-Seq profile during OIS progression displays very good correlation between SAHDs and late replicating regions (Figure 4A, S4A). Globally, replication timing was very similar across all samples. Only few regions shifted from late replicating to early replicating or vice versa (Figure 4A, Figure S4B) and the very few late
replicating regions that switch to early replication were gene poor and displayed no change in chromatin contacts (**Figure S4C-E**).

These data suggest that changes in replication timing are unlikely to play a role in SAHF formation. We then tested selected candidate proteins for their role in SAHF formation. RIF1 and LRWD1, which are known to bind late replicating regions and to affect their 3D localisation and replication timing (Foti et al., 2016; Giri et al., 2015), have no effect on SAHF formation. Likewise, depletion of Suv420H1 and Suv420H2, which deposit H4K20me3 in H3K9me3 enriched regions in OIS (Nelson et al., 2016) had no effect (**Figure 4C-E**).

One unifying feature of SAHFs is a low level of CpG DNA methylation (**Figure 4B**) and DNA methylation levels are highly correlated with replication timing in IMR90 human fibroblasts (**Figure S4F**). We therefore tested whether the maintenance DNMT1, or the de novo DNMT3A or DNMT3B DNA methyltransferases are involved in SAHF formation. We depleted these proteins at the onset of OIS formation (**Figure 4C**) and compared the results with siRNAs against firefly luciferase gene (henceforth called non-targeted or NT) as a negative control and with siRNAs against HMGA1 and HMGA2 proteins as positive controls, since depletion of these proteins is known to prevent SAHF formation (Narita et al., 2006). These experiments showed that DNMT1 knockdown prevents SAHF formation in OIS (**Figure 4D-E**), although senescence-like cell cycle arrest is maintained, as shown by cell cycle analysis following BrdU incorporation (**Figure S4G**). Instead, the knock down of DNMT3A or 3B had no significant effect. Therefore, DNMT1 is a new essential factor for SAHF formation.

**DNMT1 is associated with OIS-dependent 3D genome rewiring**

To study whether the absence of SAHFs upon depletion of DNMT1 corresponds to restoration of a normal 3D genome architecture we performed Hi-C on DNMT1 and control (NT) depleted OIS cells. The normalised Hi-C matrices from siDNMT1 treated cells display a global loss of long-range contacts when compared to control NT cells (**Figure 5A**). Hierarchical clustering of pairwise Pearson’s correlations between all Hi-C samples (at 250 kb resolution and considering contacts in the 100 Kb to 3 Mb range) revealed three main clusters, namely the proliferative condition, the OIS condition and the replicative senescent condition (**Figure S5A**). As expected, the control siRNA treated cells cluster with OIS samples. Surprisingly however, siDNMT1 treated samples cluster with the replicative senescent condition rather than with the OIS-D0 or the OIS-D6 conditions (**Figure S5A**).

Plotting the mean intra-chromosomal contact probability over distance shows an increase in contacts over the range of 500 kb to 16 Mb in the siDNMT1 samples when compared to controls, similar to RS-Senescent samples (**Figure S5B, 1C**). An increase in chromatin contacts in this range might indicate a reduced genome compartmentalization. Consistent with this hypothesis, we observed a decrease in compartmentalisation in siDNMT1 samples, which resembles the shift observed from the RS-Proliferative to the RS-Senescent state (**Figure 5B, S5C**). Furthermore, we observed a mild gain in intra SAHD interactions and a mild loss in inter-SAHFs interactions in siDNMT1 versus siNT samples (**Figure S5D**). Next, we performed 3D FISH using SAHD (A, B, and C) and non-SAHF (D) probe in siDNMT1 and siNT nuclei. Image analysis shows that the SAHD probes localizes within SAHFs in siNT samples (**Figure 5C**), as expected for OIS cells. This leads to the shortening of the distance between the most distant A and D probes in siNT (**Figure 5D**). In contrast, no SAHFs and no distance shortening were observed in siDNMT1 samples (**Figure 5D**). The A-D probe distance remains significantly shorter in siNT versus siDNMT1 even after nuclear size normalisation (**Figure S5E**). Further, DNMT1 KD prevents SAHD displacement from the nuclear periphery (**Figure
5E) in comparison to siNT samples. Finally, except for one SAHD region (probe A), the other SAHD regions (probes B and C) did not undergo decompaction in siDNMT1 (i.e. signal diameters were larger in the siNT condition compared to siDNMT1, see Figure 5F).

Together, these data show that DNMT1 plays an important role in SAHF formation and OIS-specific 3D genome rewiring, which is also consistent with the transcriptional induction of DNMT1 at early stages of OIS induction but not in RS conditions (Figure S5F). An important question is thus how does DNMT1 lead to SAHF formation. Among possible candidates, we found that gene expression of lamin B and the lamin B receptor are reduced during OIS. However, they were also reduced in RS-Senescent and DNMT1 knockdown cells, which do not form SAHFs (Figure S5G). This suggests that repression of the lamin B receptor might not be the mechanism through which DNMT1 induces SAHFs. Displacement from the nuclear periphery might be required to form SAHFs, however. To validate our observation, we performed DNA FISH on OIS-D6 cells depleted of HMGA1 and HMGA2 proteins, which are known to be required for SAHF formation (Figure 4B). Consistent with the requirement for SAHD displacement from the nuclear periphery in SAHF positive cells, FISH analysis showed the absence of displacement of a SAHD (probe A) from the periphery in HMGA depleted OIS cells (Figure S5H). We then tested whether HMGA proteins might be possible downstream mediators of DNMT1 function. The mRNA levels of HMGA1 and HMGA2 were strongly induced upon OIS, but not in RS and siDNMT1 treated cells (Figure S5I). Since HMGA2 is known to be critical for SAHF formation, these data suggest that the downregulation of HMGA2 in siDNMT1 treated cells might be the reason for the blockade of SAHF formation.

DNMT1 is known to act as a transcriptional repressor. Since its levels are induced upon OIS, the most plausible hypothesis is that DNMT1 might repress an HMGA repressor in OIS. HMGA2 is known to be repressed by the BRCA1/ZNF350/RBBP8 repressor complex (West et al., 2019). We found that these three genes were upregulated in DNMT1 depleted samples (Figure 5G, S5J). In order to analyse whether DNMT1 might downregulate BRCA1/ZNF350/RBBP8 repressor complex via DNA methylation, we performed whole genome bisulfite sequencing (WGBS) of DNMT1 and NT depleted OIS cells. Globally, mC levels in both siDNMT1 and siNT samples were well correlated, with SAHDs being hypomethylated (Figure 5H-I). However, we found that BRCA1 and ZNF350 have reduced CpG methylation levels around their promoter regions (± 3kb from TSS) (Figure 5K). In order to test whether DNMT1 might induce SAHFs via derepression of HMGA2, we overexpressed HMGA2 in OIS cells treated with DNMT1 siRNA (Figure S5K). HMGA2 overexpression rescued SAHFs (Figure 5K-L), demonstrating that HMGA2 acts downstream of DNMT1 in OIS cells. Together, these data suggest that OIS-mediated induction of DNMT1 might repress the BRCA1 and ZNF350 genes via DNA hypermethylation, leading to derepression of HMGA2 in order to induce SAHF formation.

Identification of genes associated with SAHF mediated chromatin remodelling

To analyse the effect of SAHFs on gene expression in senescence, we took advantage of multi-omics profiling to identify 3D genome architectural changes at active genes, both at the global scale as well as selectively associated to SAHF formation. First, we found that both OIS and RS cells partially lose insulation during the onset of senescence, particularly at TAD borders (Figure 6A, S6A). We then focused on active transcription start sites (TSSs), since they were shown to induce local insulation and undergo long-distance 3D interactions (Bonev et al., 2017). Surprisingly, active TSSs lose insulation in OIS samples and in control siRNA treated OIS samples, whereas the RS-Senescent and siDNMT1 treated cells maintain high insulation at active TSSs (Figure 6B). Furthermore, we found an increase in the active TSS-
TSS interactions upon OIS induction, but not in the RS-Senescent and siDNMT1 treated OIS nuclei (Figure 6C, S6B). This suggests that SAHF formation is concomitant with a global increase in interaction among active TSSs.

We reasoned that TSSs located close to SAHDs along the linear genome might be brought in spatial proximity upon SAHF formation, and that their physical interaction might reinforce transcriptional changes (Figure S6C). In order to test this hypothesis, we selected genes specifically upregulated in the presence of SAHFs (STAR Methods). Among these genes, we then further selected genes adjacent to SAHDs (ADJ) (STAR Methods). The final result was a set of 330 genes which we called SAHF-specific genes (Figure 6D, S6D, Table S4). To test whether SAHF-specific genes would have a greater chance of interacting with each other when SAHDs interact to form SAHFs, we mapped active-TSS-TSS interactions in SAHD-adjacent regions (Figure 6E). We found that SAHF-specific genes are expressed only when their TSSs interact at OIS-D4, due to SAHF formation, and both their interactions and their expression levels decrease when SAHF formation is prevented in DNMT1 knockout cells (Figure 6E). For comparison, we performed this analysis on a set of control regions, obtained by randomly shuffling SAH domains intra-chromosomally. This operation defined a set of “pseudo-SAHDs” with an interdomain distance distribution similar to that of original SAH domains. The regions adjacent to these randomly placed pseudo-SAHDs were called random adjacent SAHDs (RDM-ADJ) and genes in the RDM-ADJ regions were called “random set genes”.

Although the gene expression profiles of random set genes were matched to those of the SAHF-specific gene set (Figure 6F) and active TSS-TSS interactions in the random gene set increased during OIS, these interactions remained high in siDNMT1 treated cells, unlike for SAHD-specific genes which lost TSS-TSS interactions (compare Figure 6E with Figure 6G). This indicates that SAHF-specific genes are specifically regulated by DNMT1-dependent chromatin interactions that are mediated by SAHF formation. In order to validate TSS-TSS interactions among SAHD-specific genes in SAHF positive cells we used 3D FISH assays. As a positive set, we choose TSSs of the RHOB and CXCR gene from the SAHD specific category. As a negative control, we chose the TSS of OTOF and GPR39 genes, which are adjacent to RHOB and CXCR, respectively (Figure 6I). 3D FISH and gene expression data demonstrate that SAHD specific genes (RHOB and CXCR) become upregulated and come significantly closer than the control set of genes (OTOF and GPR39), in SAHF positive cells (Figure 6J-K, S6F-G). Together, these data indicate that SAHF-specific genes are transcriptionally upregulated upon SAHF-dependent induction of spatial proximity.

The gene ontology analysis of our OIS and RS system identifies key pathways, such as notch signalling, NF-kB pathway, DNA repair, histone modifications pathway etc, which are concordant with previously published GO analysis (Table S5) (Hernandez-Segura et al., 2017). The gene ontology analysis of SAHF-specific gene set showed an enrichment of genes for cancer, organismal injury and abnormalities and inflammatory responses. (Figure S6F). Since most of the SASP genes fall in the immune response pathways and play a vital role in senescence, we mapped their expression profiles across all samples. We found most of these genes in the ADJ region. Nevertheless, despite being present in the ADJ region we found no significant decrease in SASP gene expression upon SAHF inhibition (Figure 6H), suggesting that SASP genes are regulated independently of SAHF formation.

Finally, while our findings suggest that SAHFs are made up of SAHD heterochromatin regions, they also suggest that both transcriptionally active and inactive TSSs might be present in the vicinity of SAHFs. To test this scenario, we designed two oligopaint FISH probes spanning the TSS of two genes, one active (IGF2R) and one inactive (PNLDC1), both adjacent to SAHDs (Figure S6G). The aim of the assay was to test whether a repressed gene might locate closer to SAHFs in 3D space than a neighbouring active gene. FISH results demonstrate
that both genes co-localise in the SAHF periphery and are equidistant from the closest SAHFs (Figure S6H-L). These observations along with our previous results suggests a model in which SAHFs represent clustering of SAHDs, harbouring H3K9me3 marked silent regions, with both active and inactive genes in their vicinity. Interestingly the genes within the SAHD regions matches with genes that are downregulated in lung tissue and were upregulated in human brain tissue (Table S6), suggesting that SAHDs might be reprogrammable upon cell differentiation.

In summary, we provide here a detailed characterization of epigenome and 3D genome changes during senescence. Systematic data analysis led to the genome-wide identification of chromosomal domains that form senescence-associated heterochromatin foci upon oncogene induction. A screening of candidate factors for SAHF induction allowed us to identify DNMT1 as a novel component that induces SAHFs by stimulation of HMGA2 expression. DNMT1 depletion does not reverse the complete senescence process, however. Instead, depleted cells transition to a 3D genome conformation akin to that of cells in replicative senescence, suggesting that acute senescence induction involves SAHF formation in addition to the RS-dependent 3D genome rewiring (Figure 7).

DISCUSSION

Here, we interrogated the changes in the nuclear architecture in human fibroblasts undergoing RS and OIS at all levels of genome organisation, including RNA abundance, epigenome and 3D genome architectural changes. As fibroblasts enter senescence either via RS or OIS, individual chromosomes compact, consistent with a previous report in RS cells (Criscione et al., 2016). This compaction is not observed at all chromatin scales, however, and it rather depends on the gain of long-range contacts over tens of Mb. This gain in long range contacts might be a general feature of cells that exit mitosis, since previous work on mouse ES cell differentiation revealed a similar phenomenon in neurons (Bonev et al., 2017) and in silico polymer modelling (Figure S3M) shows a gain in long-range interactions in response to cell cycle exit and time relaxation.

A key feature that differentiates RS nuclei from OIS nuclei is SAHF formation. Although previous studies in OIS and RS have suggested that TADs and compartments change in a subtle way during OIS (Chandra et al., 2015; Criscione et al., 2016), the OIS time-course shown here identified a massive change in compartmentalization, with a decrease in RS compartmentalization and a reverse increase in OIS, driven by long-range B-compartment interactions that leads to SAHF formation. More precisely, SAHFs are formed by coalescence of conserved H3K9me3 and late replicating heterochromatin domains that we called senescence associated heterochromatin domains (SAHDs). DAPI-dense SAHF-like foci have also been observed in human, as well as in mouse neurons overexpressing GFP tagged histones (Ito et al., 2014; Sun et al., 2018; Xu et al., 2014). SAHF-like structures with gene rich regions shifting to the nuclear periphery and DAPI foci corresponding to individual chromosome territories were also observed in fibroblasts treated with chaetocin (Illner et al., 2010).

Our Hi-C 3D modeling and FISH results also highlighted a fundamental difference in SAHD organisation between RS and OIS nuclei. While the SAHDs remain partially peripheral and compacted in RS, they are displaced from the periphery and lose internal compaction in OIS. This is in agreement with a recent report demonstrating loss in compaction when heterochromatin domains move away from the periphery in mouse ES cells (Zheng et al., 2018) and with previous observations suggesting that SAHFs might arise because of release of heterochromatic lamina-associating domains from the lamina (Chandra et al., 2015; Sadaie et al., 2013; Shah et al., 2013). Our polymer modelling shows that the release of SAHDs from the
lamina is sufficient to form large foci of heterochromatin (MR scenario), consistent with a recent independent polymer model (Chiang et al., 2019). However, by accounting precisely for the large-scale, out-of-equilibrium relaxation of chromosomes and by making detailed comparisons with Hi-C and FISH data, our approach suggests in addition (MR-WI scenario) that the weakening of intra-SAHHD interactions is one key factor which, together with lamin release, quantitatively explains SAHF formation.

A key question is thus what might detach SAHDs from the lamina and decondenses SAHDs during OIS. Here, we identified DNMT1 as a key component in SAHF formation. Surprisingly, OIS cells treated with siDNMT1 have 3D structural features similar to RS cells, including a loss of SAHFs that depends on the downregulation of HMGA2, a component which was previously shown to be required for SAHF formation (Narita et al., 2006). Therefore, the present work has identified induction of DNMT1 as a requirement for SAHF formation via activation of HMGA2 expression. HMGA2 might decondense chromatin through competition with histone H1 (Catez et al., 2006; Ozturk et al., 2014) while sustaining long-range heterochromatin interactions, possibly helped via HMGA dimer formation (Frost et al., 2015). Local heterochromatin decondensation, combined with the strong reduction in lamins and Lamin B receptor expression upon induction of senescence, might be responsible for triggering loss from the nuclear periphery in OIS. This scenario is consistent with our microscopy-based observation of a strong correlation between local SAHD decondensation and SAHD detachment from the lamina.

Interestingly, SAHF formation has not only structural, but also functional consequences. Our data show that active genes adjacent to SAHDs are brought in 3D physical proximity and establish more frequent Hi-C contacts upon induction of SAHFs. Many of these genes function in senescence associated processes, such as changes in cell adhesion and the activation of cancer-related genes. Together, these data suggest that 3D genome reorganization upon exhaustion of the replicative potential or the induction of oncogene expression might be causally involved in the transcriptional responses that lead to cell senescence and protection from cancer induction. Further analysis of the rich multi-omics set of data provided here might lead to the identification of potential regulatory components involved in the induction and the biological function of senescent cells, as well as in SAHF formation upon illegitimate oncogene activation.

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AUTHOR CONTRIBUTIONS
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FIGURE LEGENDS
1. Genome organisation in OIS and RS based on distinct chromatin compartment interactions. (A) Schematic representation (DAPI staining) of the WI-38 hTERT/GFP-RAF1-ER cell-specific OIS and WI-38 primary cell-dependent RS systems. Only OIS cells show SAHF bodies (D4 onwards). Scale bar = 5 μm. (B) Normalised Hi-C contact maps for chromosome 5 at 500-Kb resolution. The lower left bottom of individual Hi-C plots represent control cells (D0 or RS-Proliferative) and upper right part displays senescence condition (OIS-D2, OIS-D4, OIS-D6, OIS-D10 or RS-Senescent). The respective eigenvectors are above the Hi-C plot. c = Control (OISD0 or RS-Proliferative), e = Treatment (OIS-D2, OIS-D4, OIS-D6, OIS-D10 or RS-Senescent). The maximum intensity for each comparison (panel) is indicated in the lower left corner. (C) Contact probability in logarithmic bins. Lines: mean values from biological replicates. (D) Representative images of chromosome territories from OIS-D0 / OIS-D6 and RS-Proliferative / senescent cells mapped via whole chromosome paint assay. Chromosome territories were delineated with Cy5 (green) labelled whole chromosome painting probes and the nuclei were counterstained with DAPI (blue). Bar = 10 μm. (E) The distributions of areas of chromosome territories are shown as box plots. Statistical significance is calculated using Mann-Whitney test. (F) Intra- and inter-compartment contact enrichment from OIS and RS samples. Data represented as bar plots showing the mean ± SD. (G) Average contact enrichment between pairs of 250-kb loci arranged by their eigenvalue (shown on top). The green bar at the bottom depicts the trend in compartmentalisation. See also Figure S1.

2. Differential intra vs inter SAHD interactions constitute SAHF
(A) Genome browser shot of differential interactions. Top: Histone modification tracks of H4 (K5ac, K8ac, K20me1, K91ac), H3 (K4me1, K4me2, K4me3, K4ac, K9me1, K9ac, K14ac, K18ac, K23ac, K27me2, K27ac, K36me3, K56ac, K79me1, K79me2) and H2 (A.Z, BK5ac, AK9ac, BK12ac, BK15ac, Bk20ac, BK120ac) modifications from IMR90 cells. Middle:
H3K9me3, mC and Repli-Seq tracks from IMR90 cells. Bottom: Differential interaction (from diffHIC) tracks from OIS-D0 vs OIS-D6 and RS-Proliferative vs RS-Senescent at 100 kb bins. The square brackets highlight the broad H3K9me3 regions gaining contacts in senescence. The scale bar represents statistically significant differential score in 100 kb bins. The blue arcs are gain in interaction while purple arc means loss. (B) Quantification of contacts within and between SAHFs in OIS and RS. Data are represented as a scatter dot plot showing the mean ± SD. (C) Schematic representation of the location of FISH probes on chromosome 5. (D) Representative 3D-DNA FISH images (z-slice) from OIS-D0 and OIS-D6 samples with A-C, B-C, AD, B-D and C-D probes. Scale bar = 5 μm. (E) Representative 3D-DNA FISH images (z-slice) from RS-Proliferative and RS-Senescent samples with A-C, B-C, AD, B-D and C-D probes. Scale bar = 5 μm. (F) Percentage of the respective FISH probes, localised within SAHF in individual nuclei, in OIS-D6 samples. (G) Quantification showing the distance of the SAHD (A, B and C) and non-SAHD (D) probes from the nuclear periphery. Data represented as box plots. Statistical significance is calculated using Mann-Whitney test. (H) Boxplot of inter-probe distances in OIS-D0 and OIS-D6. Statistical significance is calculated using Mann-Whitney test. (I) Quantification of the changes in the diameter of the individual probes in OIS-D0 and D6. Statistical significance is calculated using Mann-Whitney test. (J) Quantification showing the distances of the probe from the nuclear periphery in RS-Proliferative and RS-Senescent. Statistical significance is calculated using Mann-Whitney test. (K) Boxplot of inter-probe distances in RS-Proliferative and RS-Senescent. Statistical significance is calculated using Mann-Whitney test. (L) Quantification of the changes in the diameter of the individual probes in RS-Proliferative and RS-Senescent. Statistical significance is calculated using Mann-Whitney test. (M) Representative 3D-DNA FISH images (z-slice) from BJ rafD0 and BJ rafD6 samples with A-D probes. Scale bar = 5 μm. (N) Percentage of the respective FISH probes, localised within SAHF in individual nuclei, in BJ rafD6 samples. (O) Quantification showing the distance of the SAHD (A) and non-SAHD (D) probes from the nuclear periphery. Data represented as box plots. Statistical significance is calculated using Mann-Whitney test. (P) Boxplot of inter-probe distances in BJ rafD0 and BJ rafD6. Statistical significance is calculated using Mann-Whitney test. (Q) Quantification of the changes in the diameter of the individual probes in BJ rafD0 and BJ rafD6. Statistical significance is calculated using Mann-Whitney test. See also Figure S2 and Table S2 and Table S3.

3. Detachment from lamina partially explains 3D Genome Reorganization in OIS

(A) A two-parameter polymer model of heterochromatin organization and positioning. Each 10-kbp monomer is characterized by an epigenetic and lamina-interacting state. SAHD monomers exhibit pair-wise contact attraction and cLAD-like loci may interact preferentially with the nuclear membrane. An example of configuration evolving in a cubic box (one of its face representing the membrane) is given at the bottom right corner of the figure. Panels (B) to (D): Display the inference of polymer model for cycling/OISD0 cells. (B) Ratio between the average contact probability between SAHD regions and the total average contact probability between any pairs of loci (expected probability), as a function of the genomic distance. (C) Example of SAHD compartment. (D) Distributions of the number of small, intermediate and large, SAHF-like SAHD compartments per simulated configuration. Panels (E) to (G): Display the predictions for the time-release (TR) scenario. (E) Ratio between the observed vs expected ratio for SAHD regions at OIS-D6 and at OIS-D0. (F) Example of SAHD compartment. (G) Distributions of the number of small, intermediate and large, SAHF-like SAHD compartments per simulated configuration. Panels (H) to (J): Display the predictions for the membrane-release (MR) scenario. (H) Ratio between the observed vs expected ratio for SAHD regions at OIS-D6 and at D0. (I) Example of SAHD compartment. (J) Distributions of the number of
small, intermediate and large, SAHF-like SAHD compartments per simulated configuration. Panels (K) to (M): Display the predictions for the membrane-release with weakening of SAHD attraction (MR-WI) scenario. (K) Ratio between the observed vs expected ratio for SAHD regions at OIS-D6 and at OIS-D0. (L) Example of SAHD compartment. (M) Distributions of the number of small, intermediate and large, SAHF-like SAHD compartments per simulated configuration. (N) Distance distribution between B, C, D probes from polymer model. See also Figure S3.

4. DNMT1 knockdown prevents SAHF formation

(A) Replication-timing (RT) profiles of WI38 RAF cells at early stages of OIS. The data displayed as log2 ratios of signals from early and late S-phase fractions. Positive scale corresponds to an early replication and negative scale correspond to late replication timing. (B) Washington University Genome browser shot. Top: mC tracks from IMR90 cells. The blue colour indicates methylation levels and the grey background coverage. Data from NCBI roadmap to epigenomics. Middle: Replication timing track of un induced WI38 RAF cells (G1 phase to G2 phase) The classification of G1 to G2 phase is mentioned in materials and methods. Bottom: SAHDs falling in the displayed region. (C) Schematic overview of the knockdown experiment. OISi (oncogene induction). RNAi (administration of siRNA). Day 0 (D0) to Day 5 (D5). (D) Representative DAPI staining images (z-slice) from oncogene induced (D5), control (NT) and upon depletion of HMGA1 + HMGA2, DNMT1, DNMT3A, DNMT3B, SUV420H1, SUV420H2, LRWD1, RIF1, GAPDH. Scale bar = 10 μm. (E) Quantification of the changes in the SAHF score (details in STAR Methods) of the cells displayed in (D). Statistical significance is calculated using Mann-Whitney test. See also Figure S4.

5. DNMT1 underlie the 3D changes that differentiates RS from OIS state

(A) Normalised Hi-C contact maps shown for chromosome 5 at 500-Kb resolution. The maximum intensity for each data is indicated in the lower left corner. (B) Average contact enrichment between pairs of 250-kb loci arranged by their eigenvalue (shown on top). (C) Left panel is representative 3D-DNA FISH images (z-slice) from NT and DNMT1 depleted samples with A-C, B-C, B-D and C-D probes. Scale bar = 5 μm. Right panel display percentage of the respective FISH probes, localised within SAHFs in individual nuclei, in NT samples. (D) Boxplot of inter-probe distances in NT and DNMT1 depleted samples. Statistical significance is calculated using Mann-Whitney test. (E) Quantification showing the distance of the SAHD (A, B and C) and non-SAHD (D) probe from the nuclear periphery. Data represented as box plots. Statistical significance is calculated using Mann-Whitney test. (F) Quantification of the changes in the diameter of the individual probes in NT and DNMT1 depleted samples. Statistical significance is calculated using Mann-Whitney test. (G) BRCA1, ZFP350 and RBBP8 expression represented as the mean ± SD of two biological replicates of RNA-seq experiments for NT and DNMT1 depleted samples. (H) IGV snapshot of chromosome 10 with WGBS tracks from siNT and siDNMT1 treated OIS samples. (I) Scatter plot comparing the mC levels (percent methylation) in siNT and siDNMT1 samples. The correlation of the individual datasets is indicated on the bottom right panel. (J) IGV snapshot of the ZNF350 and BRCA2 genes, along-with WGBS tracks from NT and DNMT1 depleted OIS cells. The red box highlights the differentially methylated promoter region. (K) Quantification of the changes in the SAHF score (details in STAR Methods). Statistical significance is calculated using Mann-Whitney test. (L) An immunofluorescence experiment on OISD5 cells in DNMT1 deleted and HMGA2 overexpression condition using an antibody against HMGA2. Zoomed
cell displays a single nucleus with SAHF bodies and HMGA2 overexpression. See also Figure S5.

6. Gene expression changes associated with SAHF formation.

   (A) Average insulation score over 200kb region around TAD boundaries in OIS, RS and siRNA treated OIS cells. Lines show mean values, while dark and light shaded ribbons represent SD and 95% CI respectively. (B) Average insulation score centred on gene promoters in OIS, RS and siRNA treated OIS cells. Lines show mean values, dark and light shaded ribbons represent SD and 95% CI respectively. (C) Long range (2- to 10-Mb) inter-TAD aggregate Hi-C contact maps around pairs of active transcription start sites (TSS) in OIS-D0, OIS-D2, OIS-D4, RS-Proliferative, RS-Senescent, NT depleted and DNMT1 depleted samples within ± 40 kb. (D) Quantification of gene expression of the SAHF-specific genes following the (STAR Methods). The data displayed as box plot of log 2(FPKM +1) values. (E) Quantification of the contact strength between pairs of TSS from SAHF-specific genes represented in panel (D). Data represented as a box plot showing the mean ± SD. (F) Quantification of gene expression of the random ADJ control genes (random gene set). The data displayed as box plot of log 2(FPKM +1) values.  (G) Quantification of the contact strength between pairs of TSS from random gene set category. Data represented as a box plot showing the mean ± SD. (H) SASP gene expression represented as the mean ± SD of two biological replicates of RNA-seq experiments for NT and DNMT1 depleted OIS cells. (I) Schematic representation of the location of FISH probes on chromosome 2. (J) Representative 3D-DNA FISH images (z-slice) from OIS-D0 and OIS-D6 samples with Set 1 (RHOB and CXCR4) and Set 2 (OTOF and GPR39) probes. Scale bar = 5 μm. (K) Quantification showing the distance of the within each probe set. Data represented as box plots. Statistical significance is calculated using Mann-Whitney test. See also Figure S6.

7. Unified model of chromatin organisation in different senescence conditions.

OIS leads to DNMT1 overexpression and a simultaneous release of SAHDs from periphery. SAHDs become decompacted as they move away from periphery and gain inter-SAHD interactions to form SAHFs. Conversely, in RS conditions the SAHDs remain at the periphery with no major changes in inter or intra-SAHD interactions. However, in cells undergoing OIS, DNMT1 knockdown leads to a partial rescue of SAHDs detachment from lamin and prevention of HMGA2 overexpression. These conditions lead to the prevention of SAHF formation in DNMT1-depleted OIS nuclei and the new chromatin architecture resembles with Replicative-Senescent state.

STAR Methods

KEY RESOURCE TABLE

CONTACT FOR REAGENT AND RESOURCES SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Giacomo Cavalli (giacomo.cavalli@igh.cnrs.fr).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell Lines
WI-38hTERT/GFP-RAF1-ER and BJ-hTERT-B-RAF-V600E cells were kindly provided by Carl Mann (Carvalho et al., 2019; Jeanblanc et al., 2012). The HSF74 cells are from Lapasset et al (Lapasset et al., 2011). The cells were grown under conditions described by Jeanblanc et al. (Jeanblanc et al., 2012). Briefly, WI-38hTERT cultured in modified Eagle’s medium (Thermo-Fisher Scientific, Cat.N: 12492013), supplemented with 10% fetal bovine serum (Thermo-Fisher Scientific, Cat.N: 10270106), 1mM sodium pyruvate (Thermo-Fisher Scientific, Cat.N: 11360070), 1mM of glutamax (Thermo-Fisher Scientific, Cat.N: 35050-038), 50U of penicillin and streptomycin (Thermo-Fisher Scientific, Cat.N: 15070-063) and 0.1mM MEM non-essential amino acids (Thermo-Fisher Scientific, Cat.N: 11140-035). Cells were cultured under 5% carbon dioxide and 5% oxygen conditions. Cells become senescent in 72 hours post 4-hydroxytamoxifen induction (Sigma-Aldrich, Cat.N: H7904). The WI-38 primary cells were bought from ATCC.

METHOD DETAILS

BrdU FACS for Cell cycle analysis

Cell cycle progression was assessed by flow cytometry. Cells were pulsed 2h with 100 μM of Bromodeoxyuridine (BrdU). After trypsinization (Thermo-Fisher Scientific, Cat.N: 25200056) and wash in PBS 1% foetal bovine serum (FBS), cells were fixed in 75% ice cold ethanol. Fixed cells were washed, collected by centrifugation, treated with 2N HCl-0.5% triton X-100 30 min at room temperature, washed and neutralized in 0.1M Na2B4O7•10H2O, pH 8.5. After centrifugation, cells were incubated overnight at 4°C with primary anti-BrdU antibody (Exbio, Cat.N:11286-c100) diluted in PBS-0.5% Tween 20-1% BSA, washed in PBS 1% CFS and incubated 1h at RT with Alexa 488 labelled secondary antibody (Abcam, Cat.N: ab150113). Then, cells were washed and resuspend in PBS 1% FBS containing 50 µg/ml of Propidium Iodide (Thermo-Fisher Scientific, Cat.N: P3566) and RNase A 200µg/ml (Thermo-Fisher Scientific, Cat.N: 12091021) for 30 min at room temperature. Data acquisition were done in Gallios FACS apparat (Beckman Coulter) and analyzed using Kaluza software (Beckman Coulter). Relative cell cycle phase distributions were performed by using bivariate dot plot analysis.

siRNA transfection

siRNA transfection for image analysis was performed on 100 thousand cells. For Hi-C 1 million WI-38hTERT/GFP-RAF1-ER cells were transfected. In both cases, we use RNAiMax with a 30 nM final concentration of predesigned siRNAs form Dharmacon (product details in Key Resource Table)

Hi-C Library Preparation

The cells were dissociated using 0.25% Trypsin-EDTA (ThermoFisher, Cat.N: 25200056). After fixation for 10 min at room temperature with freshly prepared 1% formaldehyde in PBS (ThermoFisher, Cat.N: 28908), the reaction was quenched for 5 minutes by adding 0.2M glycine solution (final). Cells were permeabilized using 0.1% saponin (in PBS) and concentration was adjusted to 1x10^6 cells / mL. 1 µL/mL of fxCycle far red dye (ThermoFisher, Cat.N: F10348) and 5 µL/mL RNase A (20mg/mL) were added and samples were incubated for 30 min at room temperature protected from light with slight agitation. After washing once
with cold 1x PBS, samples were resuspended in cold 0.5% BSA in PBS at a concentration of 10^7 cells/mL and immediately processed for FACS. FACS was performed using BD FACSAria and appropriate gates were set based on the relative levels of fxCycle in order to isolate G0G1 cells. After FACS sorting, cell were pelleted and were frozen in liquid nitrogen and stored at -80°C. The RS-Senescent cells were not FACS sorted as replicative senescent cells are mostly in G1 phase. Hi-C and library preparation was carried out using the in-situ method as described previously (Rao et al., 2014) with minor modifications. In brief, cells either FACS-purified or RS-Senescence were digested overnight at 37°C using 500U of DpnII. After biotin filling, proximity ligation was carried out for 4 hours at 18°C with 2000U T4 DNA Ligase. After reverse-crosslinking, DNA was purified using ethanol precipitation and sheared to 300-400bp fragments using Covaris S220 sonicator. Ligation fragments containing biotin were immobilized on MyOne Streptavidin T1 beads (ThermoFisher Cat.N: 65602), end-repaired and a-tailed as described. NEXTflex adaptors (Bioo Scientific, Cat.N: 514101) were then ligated and fragments were PCR amplified using KAPA HiFi Library Amplification Kit (Kapa Biosystems, Cat.N: KK2620) for 8 cycles. DNA was then double-size selected using AMPure XP beads (Agencourt, Cat.N: A63881) to isolate fragments between 300 and 800bp.

**Strand-Specific Total RNA Library Preparation**

Cells were rinsed with PBS once and then lysed immediately on plate by adding TRIzol (ThermoFisher, Cat.N: 15596026) and scraped with cell scraper. The lysate was transferred to the DNase-RNase free tubes and incubated at room temperature for 5 min. Thereafter samples were vortexed for 20 seconds, 0.2x volumes of chloroform was added, tubes were mixed by inverting and samples were centrifuged at 13,000 rpm at 4°C for 15 min. The aqueous phase was then processed using the RNeasy Mini Kit (QIAGEN, Cat.N: 74104) with DNase treatment, according to the manufacturer’s instructions. Purified RNA (maximum of 1ug) was then used for library generation using the TruSeq Stranded Total RNA Library Kit according to the manufacturer’s instructions.

**ChIP and Library Preparation**

The cells were rinsed with PBS once and then fixed immediately by adding freshly prepared 1% formaldehyde in PBS (ThermoFisher, Cat.N: 28908). The reaction was then quenched by adding glycine solution (final 0.125M) for 5min. After rinsing the cells twice with 10 mL ice cold PBS. Cells were scraped thoroughly with a cell scraper (in 10 mL ice cold PBS with 1xEDTA-free Protease Inhibitors) and transfer into 50 mL tube. The cells were pelleted immediately. The pelleted cells were then frozen in liquid nitrogen and stored at -80°C until further use. We used ~ 2x10^6 cells per IP for chromatin marks. The pellets were thawed on ice, resuspended in cold cell lysis buffer (10mM Tris pH 8, 10mM NaCl, 0.2% NP-40) + 1xEDTA-free Protease Inhibitors. Cells were lysed for 30 min at 4°C, washed once with cold lysis buffer and resuspended in cold nuclei lysis buffer (50mM Tris pH8, 10mM EDTA, 1% SDS) + 1xProtease Inhibitors (20µl per 1x10^6 cells). Nuclei were lysed for 30 min at 4°C with rotation and then sonicated for 30sec on/ 30 sec off per cycle for 18 cycles using Bioruptor (Diagenode). After sonication, 10x volumes of IP dilution buffer (20mM Tris pH8, 2mM EDTA, 150mM NaCl, 1% Triton X-100, 0.01% SDS + protease inhibitors) was added, chromatin was precleared using 20µl Protein A dynabeads (ThermoFisher, Cat.N: 10002D) / 1mL for 2hrs at 4°C with rotation. Meanwhile, 25ul beads / IP were washed once with cold 0.5% BSA in PBS, and incubated with the antibody for 4-5hrs at 4°C in 0.5ml 0.5% BSA in PBS. Beads were then washed once with 0.5% BSA in PBS, added to the precleared chromatin and incubated
overnight at 4°C with rotation. Beads were then washed once with cold IP wash buffer 1 (20mM Tris pH8, 2mM EDTA, 50mM NaCl, 1% Triton X-100, 0.1% SDS), twice with high salt wash buffer (20mM Tris pH8, 2mM EDTA, 500mM NaCl, 1% Triton X-100, 0.1% SDS), once with cold IP wash buffer 2 (10mM Tris pH8, 1mM EDTA, 250mM LiCl, 1% NP40, 1% sodium deoxycholate) and twice with cold TE buffer (1mM Tris pH8, 1mM EDTA). DNA: protein complexes were then eluted twice for 30 min at 65°C in 100 µl elution buffer (100mM NaHCO₃, 1% SDS) each time. 16 µl 5M NaCl was then added and samples + inputs were reverse cross-linked at 65°C, RNase A and proteinase K treated and purified using ultrapure phenol/chloroform (ThermoFisher, Cat.N: 15593-049). Libraries were prepared using Illumina’s TruSeq ChIP Sample Preparation Kit, according to the manufacturer’s instructions with two exceptions: libraries were PCR amplified using KAPA HiFi Library Amplification Kit for 10-12 cycles to maximize complexity and the gel-based size-selection was performed after PCR amplification.

**Replication Timing Sequencing (Repli-Seq)**

Genome-wide Replication timing (RT) profiles were constructed as previously described (Marchal et al., 2018; Ryba et al., 2012). Briefly, cells were pulse labelled with BrdU and separated into early and late S-phase fractions by flow cytometry and processed by Repli-Seq. Sequencing libraries of BrdU-substituted DNA from early and late fractions were prepared by NEBNext Ultra DNA Library Prep Kit for Illumina (E7370). 50bp single end sequencing was performed on Illumina-HiSeq 2500.

**Oligopaint Probe design and synthesis**

Oligopaint libraries were constructed following the procedures described in (Beliveau et al., 2015); see the Oligopaints website (https://oligopaints.hms.harvard.edu) for further details. Libraries were ordered from CustomArray (Bothell, WA) in the 12K Oligo pool format. Genomic regions A, B, C, D used in this study were located on human chromosome 5. Hg19 coordinates, size, number and density of probes for the libraries are given in Table S3.

Gene-poor and gene-rich oligopaint libraries were discovered using the “stringent” parameter set of OligoMiner (Beliveau et al., 2015), consisting of 40-46-mer genomic sequences throughout the regions of interest. TSS oligopaint libraries were discovered using the “balanced” parameter set, consisting of 35-41-mer genomic sequences. These oligonucleotide probes can now be retrieved from the Oligopaints website (https://oligopaints.hms.harvard.edu). Each library contains a universal primer pair followed by a specific primer pair hooked to the 40-46-mer or 35-41-mer genomic sequences (124-130mers or 119-125 in total, respectively). Oligopaint libraries were produced by emulsion PCR amplification from oligonucleotide pools followed by a “two-step PCR” procedure and the lambda exonuclease method described in (Beliveau et al., 2015). The “two-step PCR” leads to secondary oligonucleotide-binding sites for signal amplification with a secondary oligonucleotide (Sec1 or Sec6) containing two additional fluorochromes, each oligonucleotide carrying three fluorochromes in total. All Oligonucleotides used for Oligopaint production were purchased from Integrated DNA Technologies (IDT, Leuven, Belgium). All oligonucleotide sequences (5’ -> 3’) used in this study are listed below (all probe details are in Table S3).

Emulsion PCR with universal primers

BB297-FWD  GACTGGTACTCGCTGACTTG
BB299-REV  GTAGGGACACCTCTGGACTGG

Two-step PCR with specific primers
- PCR1 with FWD 5' phosphorylation and REV 53mer primers
  A (red)  BB298-FWD:  /5Phos/CGTCAGTACAGGGTGATGC
  Sec6-BB187-REV:  /Sec6BS/TTGATCTTTGACCCATCGAAGC

B (green)  BB82-FWD:  /5Phos/ATCCTAGGCATACGGGAATG
  Sec1-BB281-REV:  /Sec1BS/GGACATGGGTCAGGTAGTG

B (red)  BB82-FWD:  /5Phos/GTATCGTGAAGGGAATGC
  Sec6-BB281-REV:  /Sec6BS/GGACATGGGTCAGGTAGTG

C (green)  BB81-FWD:  /5Phos/ATCCTAGGCATACGGGAATG
  Sec1-BB281-REV:  /Sec1BS/GGACATGGGTCAGGTAGTG

C (red)  BB81-FWD:  /5Phos/ATCCTAGGCATACGGGAATG
  Sec6-BB281-REV:  /Sec6BS/GGACATGGGTCAGGTAGTG

D (green)  BB293-FWD:  /5Phos/CCGAGTCTAGCGCTCTCGTCTCTCTG
  Sec1-BB294-REV:  /Sec1BS/AACAGAGCCAGCCTCTACCTG

Secondary Binding Sequences (Sec1BS and Sec6BS):
Sec1BS: CACCGACGTCGCATAGAAGAAGCGTGTG
Sec6BS: CACACGCTCTCGTCTTGCCGTGGTGATCA

PCR2 with the labeled REV 'back primer'
BB506-Alexa488  /5Alex488N/CACCGACGTCGCATAGAAGG
BB511-Cy3  /5Cy3/CACACGCTCTCGTCTTGGC
  Secondary Oligos carrying two fluorochromes
  Sec1-Alexa488-X2:
  /5Alex488N/CACACGCTCTCGTCTTGAGGTCAGTG
  /3AlexF488N/
  Sec6-Cy3-X2:
  /5Cy3/TGATCGACCAGGGCAAGAGAGCCTTGAGG
  /3Cy3Sp/

FISH

3D FISH was adapted from (Bantignies et al., 2011). Briefly, cells were fixed with 4% PFA in PBS at room temperature for 15 min. Cells were treated with PBS, 0.5% Triton for 10 minutes, washed in PBS, incubated with 0.1M HCl for 10 minutes, washed in 2XSSCT
(2XSSC, 0.1% Tween20) and incubated for 60 minutes in 50% Formamide, 2XSCCT. Probe mixture contains 20 pmol of each probe with 20 pmol of their complementary secondary, 0.8 µl of ribonuclease A (10 mg/ml), and FISH hybridization buffer [FHB; 50% formamide, 10% dextran sulfate, 2x SSC, and salmon sperm DNA (0.5 mg/ml)], in a total mixture volume of approximately 20 to 25 µl, keeping at least a 3:4 ratio of FHB/total volume. Probe mixture was added to the coverslip before sealing on a glass slide with rubber cement. Cell DNA was denatured at 80°C for 20 min, and hybridization was performed at 37°C overnight in a humid dark chamber. Cells were then washed 3 x 5 min at 37°C in 2x SSC, 3 x 5 min at 45°C in 0.1x SSC, and 2 x 5 min in PBS before DNA counterstaining with DAPI (final concentration at 0.3 mg/ml in PBS). After final washing in PBS, coverslips were mounted on slides with Vectashield (CliniSciences) and sealed with nail polish.

**Image acquisition**
Images were acquired using confocal laser scanning microscopy (Leica SP8) equipped with x63/1.4 NA Plan Apochromat oil immersion objective (pixel size of 59 nm, z-step of 300 nm). Images for SAHF screening were acquired using wide field microscopy (Zeiss Axioimager Apotome) equipped with x63/1.4 NA Plan Apochromat oil immersion objective (Pixel size of 102 nm, z-step of 240 nm).

**Whole genome bisulfite sequencing (WGBS)**
Genomic DNA was isolated from the cell using Qiagen DNeasy Blood & Tissue Kit (Qiagen, Valencia, USA) following manufacturers’ recommendations. The WGBS libraries were constructed following Illumina's recommendations using Illumina TruSeq DNA Methylation Kit and EZ DNA Methylation-Gold kit.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**ChIP-Seq Analysis**
Fastq files were aligned to the hg19 reference genome, PCR duplicates were removed using Samtools and normalized genome coverage tracks were generated from uniquely mapping reads (mapq > 30) using deepTools2. In order to avoid double counting for paired-end data, reads with mates were extended to match the fragment size defined by the two read mates, and the second read mate was discarded. Single-end reads, unmated reads, mate reads that map too far apart (> 4x fragment length) were extended by 200bp. Biological replicates were pooled and coverage was then calculated as average reads per million of mapped reads (RPM) in 10bp bins. To determine the peaks for ChIPs with narrow binding profiles (H3K4me3), datasets were uniformly processed using the MACS2 with default parameters. For broad histone marks, the peaks (or regions) we employed SICER2 with default parameters. We have called peaks in both individual replicates and in the pooled dataset. Then we used bedtools intersect to get common peaks between replicates and finally these peaks were intersected with the pooled peaks data to get the final peak coordinate and score. This method was employed on both MACS2 and SICER2 generated peaks.

**RNA-Seq Analysis**
Fastq files were aligned to the hg19 reference genome using STAR_2.4.0i in basic two Pass mode using the “Encode” options as specified in the manual. PCR duplicates were removed and RPM normalized strand-specific coverage tracks based on uniquely aligned reads were also produced using STAR in “inputAlignmentsFromBAM” runMode. Reads overlapping with annotated genes (Ensembl build hg19) were counted using the summarizeOverlaps function from the R package “GenomicAlignments” in strand-specific, paired – end mode. FPKM (Fragments per kilobase per million mapped fragments) counts and differential expression was estimated using DESeq2. Genes with FPKM ≥ 1 were considered expressed for subsequent analysis.

**Hi-C Data Analysis**

The Hi-C data analysis was performed according to Bonev et al., 2017 (Bonev et al., 2017; Olivares-Chauvet et al., 2016). Briefly, raw sequencing reads were mapped to hg19 reference genome using Bowtie2, and uniquely mapped (MAPQ > 30) reads being translated into a pair of fragment-ends (fends) by associating each read with its downstream fend. Library statistics are presented in Table S1 and include total reads sequenced and number of valid interactions after filtering (PCR duplications, no digestion etc). Downstream analysis was then performed at fragment end resolution with no binning unless otherwise stated. To generate an expected model, we shuffle the observed Hi-C contacts using the Shaman R package (https://bitbucket.org/tanaylab/shaman) with default parameters (Olivares-Chauvet et al., 2016). The trans contacts are calculated as ratio of sum of observed contacts between the regions in trans versus sum of their marginal contacts (the contacts between region of interest and all other chromosomes in trans). Related to **Figure S21**.

**Contact Probability**

Contact probability as a function of the genomic distance was calculated as previously described (Bonev et al., 2017; Olivares-Chauvet et al., 2016). We calculated the distribution of the Hi-C contacts either as a log10 contact probability in log10 genomic distance bins, or in order to better visualize differences between conditions, as a simple contact probability (sum of the observed counts per log2 bin, divided by the all of the observed contacts, without normalizing for the bin size). We measured the “contact probability scaling” exponent as the slope of the best-fit line of the cis-decay curve when plotted on log-log axes, within a chosen range of distances. In addition to the genome-wide contact probability curves which were aggregated across individual chromosomes, we also generated contact probability maps where we considered only contacts that are located completely within a TAD (intra-TAD cis-decay curve) or inter-TAD. For **Figure 1F** we also took into consideration whether the interaction is across compartments (A-B or B-A) or within a compartment (A-A, B-B).

**Insulation, TAD, and TAD Boundary Calling**

Calculation of insulation scores and boundary calling was performed as described (Bonev et al., 2017; Nagano et al., 2017; Olivares-Chauvet et al., 2016). Briefly insulation was based on observed contacts the definition of insulation score was taken from Nagano et al. and Olivares-Chauvet et al. (Nagano et al., 2017; Olivares-Chauvet et al., 2016). The insulation score was computed individually on replicates and on the pooled contact map at 1Kb resolution within a region of ± 250Kb and is multiplied by (-1) so that high insulation score represents strong insulation. To account for any genome-wide changes in the insulation score, we further
normalized it by multiplying with a factor defined as the average insulation score across all 1Kb genomic bins in each cell type, divided by the mean of all cell types. These insulation scores were termed as normalised insulation scores. Domain boundaries were then defined as the local 2Kb maxima in regions, where the insulation score is above the 90% quantile of the genome-wide distribution. Boundaries within 10Kb of each other were merged and only boundaries, which were detected in at least 2 biological replicates, were retained. Domains shorter than 50Kb were discarded. Related to Figure 6.

**Identification of A and B Compartments**

TADs were assigned to either the A or the B compartment as described (Bonev et al., 2017; Nagano et al., 2017). In brief, we used k-means clustering (K = 2) on the log2 ratio of observed and expected (based on genomic length) inter-TAD contacts in trans. Domains in each cluster exhibit distinct signature based on the enrichment of replication timing (Figure S1G), prompting us to assign cluster 1 to the B, inactive compartment and cluster 2 to the A, active compartment. To determine the compartment strength, we calculated the log2 ratio of observed versus expected contacts (intrachromosomal separated by at least 10Mb) either between domains of the same (A-A, B-B) or different type (A-B). As another measure of compartmentalization which relies on the intra-chromosomal contacts, we calculated the dominant eigenvector of the contact matrices binned at 100Kb as described (Lieberman-Aiden et al., 2009) using scripts available at (https://github.com/dekkerlab/cworld-dekker). Related to Figure 1, 5).

**Image Analysis**

3D-FISH analysis was performed using Imaris software and its XT module. After Background substraction and Gaussian filter (sigma = 1), distances between FISH probes were calculated between the nearest mutual neighbour centers of mass of segmented FISH. For distance to nuclear periphery, nuclei were segmented in 3D using DAPI staining and the distances between FISH centers of mass and closest DAPI segmentation contour were calculated. SAHF bodies were segmented using DAPI staining, and FISH centers of mass present within SAHF bodies were considered as colocalized. Distances between FISH spots and SAHF bodies were calculated between FISH and SAHF centers of mass. Diameters of FISH signals were calculated using ImageJ software from Gaussian filtered (sigma = 1) and maximum projected images, using the Full Width at Half Maximum (FWHM) of Gaussian fitted curves to the intensity measured along scan-lines passing through FISH loci. Areas of chromosome paints and SAHF scores were calculated using MATLAB (Related to Figure 2). For chromosome paints areas, channels were smoothed using Gaussian filters (sigma = 3 for DAPI; sigma = 1 for chromosome paints) and segmented in 2D; only nuclei displaying 2 chromosome paints segmented objects were considered for analysis (Related to Figure 1).

For SAHF score quantification, images were first maximum projected along the z-axis. Nuclei were then segmented from Gaussian filtered (sigma = 3) DAPI staining. SAHF score was calculated using the coefficient of variation (CV) of DAPI intensity (intensity values smoothed using a Gaussian filter with sigma = 1). Briefly, the standard deviation and the mean DAPI intensity of each nucleus were calculated. CV were then calculated as the ratio of the standard deviation over the mean intensity (Related to Figure 4-5) (Contrepois et al., 2012).
**SAHF specific gene expression analysis**

To identify genes that might be regulated by the formation of SAHFs, we selected the genes that i) do not show significant changes in transcription at OISD2 (OISD2 vs OISD0, padj > 0.05), ii) are significantly up-regulated upon SAHF formation at OISD4 (OISD4 vs OISD0, log2FC > 1, padj <=0.05). Then, the subset of these genes that either overlap with SAHD domains or are located at +/-3Mb from the closest SAHD (Adjacent SAHD: ADJ) were selected and their transcription was quantified. This genes set was termed as SAHF-Specific genes. The GEO was performed using DAVID Bioinformatics resources (Huang da et al., 2009a, b). Related to Figure 5.

**Polymer Modelling**

We modelled the genomic region from 100 to 180.94 Mbp of chromosome 5 by a semi-flexible self-avoiding block copolymer, consisting of 8094 beads of 10kbp, each of size b, moving within a face-centered cubic lattice of size SxSxS (Olarte-Plata et al., 2016). We used rigid wall condition to mimic the confinement of the region inside a chromosome territory, one face of the cubic box (z=0) representing the nuclear membrane (Figure 3A).

For the region of interest, we collected the position of the SAHDs (Table S2) and of the constitutive LADs (Lenain et al., 2017). To each 10kbp bead i, we assigned a state \((e_i, l_i)\) regarding its epigenetic \((e_i)\) and lamina-interacting \((l_i)\) status: \(e_i=1\) if the bead is inside a SAHD (=0 otherwise), and \(l_i=1\) if the bead is inside a cLAD (=0 otherwise). The total energy of a given configuration of the polymer is then given by:

\[
H = \frac{k}{2} \sum_{i=1}^{N=1} (1 - \cos \theta_i) + \varepsilon \sum_{i,j} \delta e_i e_j \delta_{i,j} + \gamma \sum_i l_i \Delta_i
\]

With k the bending rigidity of the polymer, \(\theta_i\) the angle between the bond vectors \(i\) and \(i+1\), \(\varepsilon(<0)\) the contact energy between SAHD beads, \(\delta(i,j)=1\) if beads \(i\) and \(j\) occupy nearest-neighbour sites on the lattice (=0 otherwise), \(\gamma(<0)\) the contact energy between a cLAD bead and the membrane, and \(\Delta_i=1\) if bead \(i\) occupy a site at the membrane (=0 otherwise). \(\varepsilon\) accounts for the capacity of proteins bound to heterochromatic regions like HP1 to oligomerize (Canzio et al., 2013; Larson et al., 2017; Strom et al., 2017), while \(\gamma\) reflects still unknown molecular mechanism that lead to the preferential location of cLAD at the nuclear periphery. The dynamics of the chain follows a simple kinetic Monte-Carlo scheme with local moves using a Metropolis criterion applied to \(H\), as described in (Ghosh and Jost, 2018). The values of \(k(=2.25kT)\), \(b(=95 nm)\) and \(S(=2\mu m)\) were fixed using the coarse-graining strategy developed in (Ghosh and Jost, 2018) for a 10-nm fiber model and a volumic density \(\rho=0.01 bp/nm^3\) (estimated by dividing the size of a diploid human genome 6 Gbp by the typical volume of cycling WI-38 nuclei (600 \(\mu m^3\)).

Time mapping between the simulation and experimental time were roughly performed by considering the dynamics of relaxation of the long-range contacts between OISD0 and OISD6 (Figure S3M). Such mapping leads to a time-evolution of the mean squared displacement \(MSD(t)(in \mu m^2)\approx 0.003 t^{1/2}\) with time \(t\) given in seconds, in agreement with typical experimental estimation of MSD in human cells (Lucas et al., 2014). Considering the simplicity of the model, estimated times should be viewed as orders of magnitude of real times. For a
given parameter set, we simulate 200 independent trajectories starting from random, compact, knot-free initial configurations as in (Ghosh and Jost, 2018).

We estimated $\epsilon$ and $\gamma$ parameters for OISD0 cells by comparing model predictions with various experimental data. $\epsilon=$0.055 $kT$ was estimated by adjusting the predicted evolution of the “observed vs expected” ratio $P_e(s)/P_{tot}(s)$ as a function of the genomic distance $s$ to the corresponding experimental values (~2.2 fold), with $P_e(s)$ the average contact probability between SAHD regions and $P_{tot}(s)$ the total average contact probability between any pairs of loci, sometimes called the expected contact probability (Figure 3B). $\gamma$$\sim$0.6 $kT$ was estimated by assuming that the median distance to nuclear periphery of LAD monomers was ~700nm (see the position of B & C probes that are in LADs in Figure 2G). Interestingly, the resulting LAD organization is fully consistent with independent experimental measurements of LAD positioning inside mammalian cells performed by Kind et al (Kind et al., 2013) (Figure S3B).

In particular, we retrieved that ~30% of the LADs remains close to the periphery in every cell but that the identity of the regions bound to the membrane is very stochastic and changes at every cell cycle.

For a given polymer configuration, dense SAHD compartments (Figure 3 & S3) were determined by

1. convoluting the position of each SAHD monomer on the lattice by a 3D Gaussian of width $\lambda=270$ nm to create a 3D matrix of intensity $M(i,j,k) = \sum_{(u,v,w)} \Theta(u,v,w) \exp(-2[(i-u)^2 + (j-v)^2 + (k-w)^2]/(\lambda^2))$ is occupied by a SAHD bead;

2. by thresholding this matrix to half its maximal value to focus on dense clusters ($M\leftarrow(M\geq30)$)

**TADbit Models**

We binned at 200 kb the normalized Hi-C matrices of chromosome 5. These matrices of interactions were further normalized through Z-score transformation, and used as input to establish the restraints of the modeling. The modeling parameters of max_dist, up_freq and low_freq were optimized as described (Bau and Marti-Renom, 2012). From this optimization step we selected the set of parameters that globally maximized the satisfaction of restraints in all the ensemble of time points considered (max_dist=3000, up_freq=1.0, low_freq=-1.0). We then generated four thousand models from which we kept the best one thousand for the analysis. Each of these one thousand models is meant to represent the chromatin conformation of a subpopulation of cells within the total pull used in the Hi-C experiment. The general methodology follows each of the steps defined in (Bau and Marti-Renom, 2012; Serra et al., 2017) and was automatized using TADbit (Serra et al., 2017). The 3D models were finally scaled to match sizes observed by FISH. To do so, we computed the median of the distances between the A, B, C, and D coordinates in the OIS sample 3D models and compared it with the median of the same distances observed by FISH. We used the ratio of these medians to correct the sizes of the 3D models (they were overall 3.72 times larger). Distances were extracted independently in each of the one thousand model of each time point. Finally, to visually inspect the subpopulations of models generated at each time point, we clustered them by structural similarity. Each cluster of 3D models representing a major conformational state of the chromatin within the subpopulation of cells. The clustered models were then uploaded to TADkit (https://github.com/3DGenomes/TADkit) in the MuG VRE (http://multiscalenomics.eu) for visual inspection and alignment with other genomic features, and chromatin marks. Related to Figure 2-3.
WGBS Analysis

Raw reads were trimmed using trim galore. The reads that passed trim galore were analysed using methpipe package (version 3.4.3). Briefly the reads were mapped to hg19 genome reference using “walt”, and PCR duplicates were removed with “duplicate-remover” in methpipe package. The output files were then analyzed using “methcounts” to obtain methylation levels at each cytosine site for both strands. Then “merge-methcounts” was used to merge all replicates. Since CpG existence is symmetrical in nature we employed “symmetric-cpgs” to produce a single value for each CpG pair. CpG sites which were covered with at least 10 reads were taken for downstream analysis. Library statistics are presented in Table S1 and include total reads sequenced. These tracks were then used in Figure 5I, J.

Repli-Seq Analysis

Reads of quality scores above 30 were mapped to hg19 reference genome using bowtie2. Approximately 7 million uniquely mapped reads were obtained from each library. Read counts were binned into 5kb non-overlapping windows, and log2 ratios of read-counts between early and late fractions were calculated. For classifying the replication timing into different replication timing regions, we follow the following cutoff on the Log2 ratio between early and late fractions. The following definition taken in Figure 4B

G1 phase: Log 2 ratio from 1 and above
S1 phase: Log 2 ratio from 0.5 to 1
S2 phase: Log 2 ratio from 0 to 0.5
S3 phase: Log 2 ratio from -0.5 to 0
S4 phase: Log 2 ratio from -1 to -0.5
G2 phase: Log 2 ratio from -1 and below

DATA AND SOFTWARE AVAILABILITY

Software

The R package to compute the expected tracks and the Hi-C scores is freely available at: https://bitbucket.org/tanaylab/shaman. Processing of the raw-reads is done using https://bitbucket.org/tanaylab/schic2 as described above.

Data Resources

The accession number for the data reported in this paper is GEO: GSE130306
The 3D model of OIS and RS are available at https://dev.multiscalegenomics.eu/data_repositories/data_senescence.php

SUPPLEMENTAL INFORMATION
Supplementary Figures:

S1. Global changes in chromatin organisation during senescence, Related to Figure 1.

(A) Representative images from the Senescence-Associated β-galactosidase staining in OIS-D0, OIS-D2, OIS-D4, OIS-D6, RS-Proliferative and RS-Senescent. Scale bar = 100 μm. (B) Percentages of cells stained positive for Senescence-Associated β-galactosidase staining. Data represented as bar plot showing mean ± SD (n = 3). (C) Cell cycle profiles from FACS analyses of pulsed BrdU incorporation versus propidium iodide fluorescence (DNA content) in OIS-D0, OIS-D2, OIS-D4, OIS-D6, RS-Proliferative and RS-Senescent. (D) A representative image of giemsa staining performed on un-induced WI-38 hTERT/GFP-RAF1-ER and WI-38 primary cells showing a normal Karyotype. (E) FACS sorting for cells in G0G1+ phase of the cell cycle in OIS system and for WI-38 primary cells. The upper panel display FACS profile and bottom panel the sorted population. (F) Log-log contact probability as a function of the genomic distance. The exponent γ represents the mean slope ± SD of the best-fit line between 100Kb and 2Mb. (G) Representative images from the Senescence-Associated β-galactosidase staining in BJ raf D0, BJ raf D6, HSF74-Proliferative and HSF74-Senescent cells. Scale bar = 100 μm. (H) Percentages of cells stained positive for Senescence-Associated β-galactosidase staining. Data represented as bar plot showing mean ± SD (n = 3). (I) Cell cycle profiles from FACS analyses of pulsed BrdU incorporation versus propidium iodide fluorescence (DNA content) in BJ raf D0, BJ raf D6, HSF74-Proliferative, HSF74-Senescent (J) Schematic representation (DAPI staining) of the BJ-D0, BJ-D6, HSF74-Proliferative, HSF74-Senescent and HSF92 cells. Only BJ-D6 cells show SAHF bodies. Scale bar = 5 μm. (K) Normalised Hi-C contact maps for chromosome 5 at 500-Kb resolution. For all maps the lower left bottom of individual Hi-C plots represent control cells (D0 or RS-Proliferative) and upper right part displays senescence condition (OIS or RS-Senescent). The maximum intensity for each comparison (panel) is indicated in the lower left corner. (L) Contact probability in logarithmic bins. Lines: mean values from biological replicates. (M) Enrichment for replication timing in the two compartments. (N) Pearson correlation matrix displaying the correlation [from -1 (blue) to +1 (red)] between the intra-chromosomal interaction profiles along chromosome 5. The lower left bottom of individual Hi-C correlation matrix represent control cells (D0 or RS-Proliferative) and upper right part displays senescence condition (D6 or RS-Senescent). The eigenvector are above the Hi-C matrix.

S2. Identification and validation of SAHDs, Related to Figure 2.

(A) IGV genome browser snapshot of the ChIP-Seq profiles of OIS-D0 and OIS-D6 cells displaying the conservation of H3K9me3 and H3K27me3 domains during OIS. (B) Average enrichment of H3K9me3 modification in SAHD and control regions in D0, D6, RS-Proliferative and RS-Senescent. (C) Enrichment of replication timing regions (early replicating - G1, S1, S2, S3, S4, very late replicating -G2) across SAHD and Control regions. (D) Enrichment of different classes of LADs (cLADs, fLADs, ciLADs, fiLADs) across SAHD and Control regions. The regions were taken from Lenain et al. (Lenain et al., 2017). (E) Enrichment of different classes of isochores (L1, L2, H1, H2, H3) across SAHD and Control regions. (F) Enrichment of major repeat classes (LINE, SINE, LTR, Simple repeats) across SAHD and Control regions. (G) Enrichment of Ref-Seq genes across SAHD and Control regions. (H) Representative SAHD regions displaying the ChIP tracks and the observed / expected profiles of the respective datasets. (I) Trans-inter-SAHD interactions from OIS and RS samples. Data represented as bar plots showing the mean ± SD. (J) Oligopaint probe design. View of full chromosome 5 - p arm interactions in OIS D0, D6, RS-Proliferative and RS-Senescent maps along with the ChIP profiles. The regions selected for FISH probe designing...
are highlighted in boxes. (K) Boxplot of inter-probe distances form Figure 2H and 2K, normalised by mean volume in OIS and RS system. Statistical significance is calculated using Mann-Whitney test. (L) IGV genome browser snapshot of the Repli-Seq and ChIP-Seq profiles of BJ cells from ENCODE project and Becker et al (Becker et al., 2017) respectively, along with OIS-D0 and OIS-D6 H3K9me3 profiles in chromosome 10. (M) Quantification of contacts within and between SAHDs in BJ-D0 and BJ-D6 condition. Data showing the mean ± SD. (N) Snapshot of the MuG-VRE displaying, right panel- 3D model from chromosome 5 at 100 kb resolution of OIS-D6 condition, left panel - Hi-C matrix, and bottom panel – ChIP tracks from OIS-D6 and SAHD tracks. The red line in the 3D model highlights the two SAHD positions (A and D), which are also highlighted in the Hi-C matrix. (O) Boxplot of inter-probe distances form calculated from the TADbit 3D models in OIS-D0 and OIS-D6.

S3. In silico polymer modelling of OIS conditions, Related to Figure 3.

Panels (A) to (C): Inference of a model for cycling/D0 cells. (A) Experimental and simulated maps of the ratio between the observed and expected probability. (B) Cumulative probability to find a LAD in a fraction of the total volume measured starting from the membrane. Experimental data were taken from Kind et al (Kind et al., 2013). (C) Typical snapshot extracted from simulation. We separated cLAD and SAHD monomers and isolated dense SAHD compartments.

Panels (D) to (F): Predictions for the time-release (TR) scenario. (D) Experimental and simulated maps of the ratio between the observed and expected probability. (E) Probability distribution function to find a SAHD monomer at a given distance to the membrane. (F) Typical snapshot extracted from simulation. We separated cLAD and SAHD monomers and isolated dense SAHD compartments.

Panels (G) to (I): Predictions for the membrane-release (MR) scenario. (G) Experimental and simulated maps of the ratio between the observed and expected probability. (H) Probability distribution function to find a SAHD monomer at a given distance to the membrane. (I) Typical snapshot extracted from simulation. We separated cLAD and SAHD monomers and isolated dense SAHD compartments.

Panels (J) to (L): Predictions for the membrane-release with weakening of heterochromatin attraction (MR-WI) scenario. (J) Experimental and simulated maps of the ratio between the observed and expected probability. (K) Probability distribution function to find a heterochromatin monomer at a given distance to the membrane. (L) Typical snapshot extracted from simulation. We separated cLAD and SAHD monomers and isolated dense SAHD compartments.

(M) Generic long-range relaxation of chromosome. Ratio between the total average contact probability between any pairs of loci (expected probability) at a given time upon senescence entry and the expected probability for D0 cells as a function of the genomic distance, normalized by the corresponding value at 1Mbp. (N) The correlation between Polymer model (Y axis) and TADbit models (X axis). The X and the Y axes represent distances computed from respective models, between all 100 kb bins, over 80 Mb region on chromosome 5 (100 to 180 Mb). (O) The correlation between Polymer model (Y axis) and TADbit models (X axis). The X and the Y axes represent distances computed from respective models, between all SAHDs lying over the modelled 80 Mb region of chromosome 5 (100 to 180 Mb).
S4. Replication timing and methyl cytosine levels associated with SAHDs, Related to Figure 4.

(A) Schematic overview of the Repli-Seq experiment. Cells pulse labelled with BrdU and sorted into early and late S-phase followed by immunoprecipitation and sequencing. (B) Correlation of genome-wide RT profiles of WI38 RAF cells at early stages of OIS. (C) The differential RT regions (grey boxes 1-5) identified by unsupervised clustering of RT-variable regions. The heat map shows the RT ratios \[= \log_2(\text{early/late})\]. (D) Representative examples of RT profile alterations among different clusters in OIS. The positive values correspond to early replication and negative values to late replication. The coloured box highlights the differential RT region. (E) Boxplot displaying number of genes within each individual differential RT cluster. (F) Scatter plot comparing the RT profiles and mC levels (percent methylation) in IMR90, K562 and H1 cells. The RT profiles and mC profiles for IMR90, K562 and H1 cells were downloaded from ENCODE and NCBI Roadmap to epigenomics project. The correlation of the individual datasets is indicated on the bottom right panel. (G) Cell cycle profiles from FACS analyses of pulsed BrdU incorporation versus propidium iodide fluorescence (DNA content) in siSNMT1 and NT treated cells.

S5. Changes in chromatin architecture in DNMT1 depleted cells under OIS conditions, Related to Figure 5.

(A) Pairwise Pearson’s correlation between Hi-C samples (at 100Kb resolution and considering only contacts separated by at least 100Kb and not more than 2.6Mb). (B) Contact probability over distance. Lines: mean values from biological replicates. (C) Intra and Inter compartment contact enrichment from OIS and RS samples. Data represented as bar plot showing the mean ± SD. (D) Quantification of contacts within and between SAHDs in siNT and siDNMT1 treated cells. (E) Boxplot of inter-probe distances form Figure 5D, normalised by mean volume. Statistical significance is calculated using Mann-Whitney test. (F) DNMT1 expression represented as the mean ± SD of three biological replicate RNA-seq experiments for RS-Proliferative, RS-Senescent samples, OIS-D0, OIS-D2 and OIS-D4 samples. For OIS-D6, OIS-D10, NT depleted and DNMT1 depleted samples, expression represented as the mean ± SD of two biological replicate RNA-seq experiments. (G) LMBB1, LMNB2 and LBR expression represented as the mean ± SD of three biological replicate RNA-seq experiments for RS-Proliferative, RS-Senescent, OIS-D0, OIS-D2 and OIS-D4 samples. For OIS-D6, OIS-D10, NT depleted and DNMT1 depleted samples, expression represented as the mean ± SD of two biological replicate RNA-seq experiments. (H) Quantification showing the distance of the SAHD - A and non-SAHD - D probe from the nuclear periphery. Data represented as box plots. Statistical significance is calculated using Mann-Whitney test. (I) HMGA1 and HMGA2 expression represented as the mean ± SD of three biological replicate RNA-seq experiments for OIS-D0, OIS-D2, OIS-D4, RS-Proliferative and RS-Senescent samples. For D6, D10, NT and DNMT1 expression represented as the mean ± SD of two biological replicate RNA-seq experiments. (J) Relative fold change expression of BRCA2, ZNF350 and RBBP8 gene normalised to GAPDH expression levels. The data represents the mean and SD from 3 independent replicates. (K) Schematic overview of the HMGA2 rescue experiment. OISi (oncogene induction), RNAi (administration of siRNA). 2 days before oncogene induction (D -2), Day 0 (OIS-D0) to Day 5 (OIS-D5). HMGA2 over expression (O/E) at 2 days before ois induction ((D -2)).
S6. SAHF formation leading to TSS-TSS interaction in a specific subset of genes, Related to Figure 6.

(A) Global insulation represented as box plot in OIS, RS, NT and DNMT1 depleted samples. (B) Quantification of the contact strength between pairs of inter-TAD active TSS. Data are represented as a scatter dot plot showing the mean ± SD. (C) Schematic representation of the SAHD specific gene expression analysis. The linear view of genome displays a representative view of the SAHDS and other regions including the SAHD adjacent regions (ADJ). After oncogene induction, the genome being organised into SAHFs, with SAHDS aggregated in the centre. (D) The overlap between genes upregulated in OIS-D4 (OIS-D4 vs OIS-D0) and genes that fall in ADJ region. (E) CXCR and RHOB expression represented as the mean ± SD of three biological replicate RNA-seq experiments for OIS-D0 and two replicate RNA-seq experiments for OIS-D6. (F) Gene ontology analysis of genes in SAHD specific ADJ genes. (G) Schematic representation of the location of FISH probes on chromosome 6. (H) Representative 3D-DNA FISH images (z-slice) from OIS-D6 samples with IGF2R and PNLDC1 probes. Scale bar = 5 μm. (I) Quantification showing the distance of the within the FISH probe sets and between the DAPI marked SAHF body and FISH probes. Data represented as box plots. Statistical significance is calculated using Mann-Whitney test. (J) IGF2R and PNLDC1 expression represented as the mean ± SD of three biological replicate RNA-seq experiments for OIS-D0 and two replicate RNA-seq experiments for OIS-D6.

Supplemental Tables

Table S1. Summary Statistics for the Dataset Generated in this Study (STAR Methods).

Table S2. Genomic coordinated of SAHD regions (mapped on hg19 assembly), Related to Figure 2.

Table S3. Libraries for Oligopaint probes. Coordinates, genomic sizes, number of probes and coverage of the Oligopaint probes. Related to Figure 2.

Table S4. List of SAHF specific genes and Random set genes. Related to Figure 6.

Table S5. List of differentially expressed genes and pathways in OIS and RS samples.

Table S6. GO of genes present in SAHD regions.

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Figure 1

A

OIS-D0  OIS-D2  OIS-D4  OIS-D6  OIS-D10

OIS

RS

B

Chromosome 5

OIS-D2  OIS-D4  OIS-D6  OIS-D10

RS-Proliferative  RS-Senescent

C

D

E

F

G

Figure 1
1 monomer = 10kbp

attraction $\varepsilon$ between SAHD monomers

interaction $\gamma$ between LAD monomers and periphery

neutral monomer

LAD

SAHD

LAD+SAHD

nuclear membrane

$\text{D0/cycling}$

$\text{TR scenario}$

$\text{MR scenario}$

$\text{MR- WI scenario}$

Figure 3
Figure 4

A

Chromosome 1 (mb)

RT Log2 Ratio

OIS_Ohrs_Rep1

OIS_Ohrs_Rep2

OIS_12hrs_Rep1

OIS_12hrs_Rep2

OIS_24hrs_Rep1

OIS_24hrs_Rep2

OIS_36hrs_Rep2

Chromosome 2 (mb)

RT Log2 Ratio

OIS_Ohrs_Rep1

OIS_Ohrs_Rep2

OIS_12hrs_Rep1

OIS_12hrs_Rep2

OIS_24hrs_Rep1

OIS_24hrs_Rep2

OIS_36hrs_Rep2

B

Chr 5 2 mb

45 mb

mC

G1

S1

S2

S3

S4

G2

SAHD

C

RNAi RNAi RNAi

SAHF Quantification

D

OIS D5 NT HMGA1_2 DNMT1 DNMT3A DNMT3B

SUV420H1 SUV420H2 LWRD1 RIF1 GAPDH

E

SAHF score

p < 0.0001
Figure 5

A

Chromosome 8

OIS-D0   OIS-D4   siNT

RS-Proliferative   RS-Senescent

B

OIS-D0   RS-Proliferative   siNT

RS-Senescent   siDNMT1

C

siNT   siDNMT

D

p = 0.002   p = 0.03   ns   ns

E

p = 0.02   p = 0.04   p = 0.002   ns

F

ns   p = 0.03   p = 0.004   ns

G

siDNMT1   siNT

RBBP8   BRCA1   ZNF350

H

Chr 10

siDNMT1   siNT   SAHDs

I

Global   Non-SAHD

SAHD

J

BRCA1   ZNF350   RBBP8

siDNMT1   siNT

K

p < 0.001   p < 0.001

L

DAPI   HMGA2   Merge
Figure 6

A

B

C

D

E

F

G

H

I

J

K

Figure 6
Partial detachment of SAHDs from Lamina
Weak Compartmentalisation
SAHD compaction

Gain of Intra-SAHD Interactions

Loss of Intra-SAHD Interactions

Gain in Inter-SAHD Interactions

Detachment of SAHD from Lamina
Strong Compartmentalisation
SAHD decompaction

Chromosome territory
SAHDs
Adjacent (ADJ) chromatin domains
Distal chromatin domains
Lamina

Up-regulation
Down-regulation
Knockdown mode
Interaction frequency

No SAHF

DNMT1 Knockdown

HMGA2

DNMT1

Figure 7
REAGENT or RESOURCE

Antibodies
H3K4me3, rabbit, monoclonal
H3K9me3, rabbit, polyclonal
H3K27me3, rabbit,
Anti 5-bromodeoxyuridine, mouse, monoclonal
Anti-HMGA2

Chemicals, Peptides, and Recombinant Proteins
siRNA-DNMT1
siRNA-DNMT3A
siRNA-DNMT3B
siRNA-NT4
siRNA-HMGA1
siRNA-HMGA2
siRNA-SUV420H1
siRNA-SUV420H1
siRNA-SUV420H2
siRNA-LRWD1
siRNA-LRWD1
siRNA-RIF1
siRNA-GAPDH
Lipofectamine RNAiMAX
Cell Line Nucleofector kit R
4-Hydroxy-Tamoxifen
Senescence Beta Galactosidase Staining Kit

Critical Commercial Assays
FxCycle, Far Red
TruSeq ChIP Sample Preparation Kit
TruSeq Stranded Total RNA Library Kit
NEBNext Ultra DNA Library Prep Kit for Illumina
TruSeq DNA Methylation Kit

Deposited Data
Raw Hi-C data generated from WI38 and BJ OIS system
Raw Hi-C data generated from RS system, HSF74 and HSF92 cells
Raw Hi-C data generated from DNMT1 and scramble siRNA treated samples
H3K4me3 ChIP-Seq in OISD0, OISD2, OISD6 and OISD10 cells
H3K4me3 ChIP-Seq in RS-Proliferative and RS- Senescent cells
H3K9me3 ChIP-Seq in OISD0, OISD2, OISD6 and OISD10 cells
H3K9me3 ChIP-Seq in RS-Proliferative and RS- Senescent cells
H3K27me3 ChIP-Seq in OISD0, OISD2, OISD6 and OISD10 cells
H3K27me3 ChIP-Seq in RS-Proliferative and RS- Senescent cells
Strand Specific total RNASeq from OISD0, OISD2, OISD4, OISD6 and OISD10 cells
Strand Specific total RNASeq from RS-Proliferative and RS- Senescent cells
WGBS DNMT1 depleted OIS cells
WGBS NT depleted OIS cells
Replication timing from OISD0 cells
Replication timing from oncogene induced raf cells at 12 hour, 24 hour and 36 hour

Datasets Reanalyzed
WGBS IMR90
WGBS H1
WGBS K562
Hi-C data HUVEC Proliferative Cells
Hi-C data IMR90 Proliferative Cells
Hi-C data MSC Proliferative Cells
Hi-C data HUVEC Proliferative Cells
Hi-C data IMR90 Proliferative Cells
Hi-C data MSC Proliferative Cells
Hi-C data WI38 raf Proliferative Cells
Hi-C data WI38 raf oncogene senescent Cells
BJ H3K9me3 ChIP-Seq
BJ Repli-Seq data

Experimental Models: cell Lines
WI-38hTERT/GFP-RAF1-ER
WI38
BJ-hTERT-B-RAF-V600E
Skin Fibroblasts from 74 year old Human
Skin Fibroblasts from 92 year old Human

Software and Algorithms
Shaman R package for a-parametric Hi-C normalization
imageJ
Bowtie2 v2.2.0
Samtools v0.1.19
DeepTools2
CSAW
STAR v2.5
DESeq2
MACS2
SICER2
methpipe
C-world (Hi-C analysis software)
GraphPad Prism
MuG VRE
methpipe
SOURCE

Millipore
Abcam
Active Motif
Exbio
Abcam

GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
GE Healthcare Dharmacon
ThermoFisher Scientific
Lonza Bioscience
Sigma-Aldrich
Cell Signaling Technology

ThermoFisher
Illumina
Illumina
NEB
Illumina

This study
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UCSD Human reference Epigenome Mapping Project
UCSD Human reference Epigenome Mapping Project
ENCODE
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(Jeanblanc et al., 2012)

(Carvalho et al., 2019)

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N/A
N/A
N/A
N/A
Andrew D Smith
Job Dekker lab
N/A
N/A
N/A
IDENTIFIER

Cat# 04-745; RRID: AB_1163444
Cat# ab8898; RRID: AB_306848
Cat# 39155; RRID: AB_2561020
Cat# 11-286-C100; RRID: AB_10732986
Cat# ab97276; RRID: AB_10679322

M-004605-01-0010
M-006672-03-0005
M-006395-01-0005
D-0012-10-04-20
MQ-004597-02-0010
M-01349-5-02-0010
D-013366-01-0002
D-013366-02-0002
D-018622-23-0002
D-016934-01-0002
D-016934-02-0002
MQ-027983-01-0002
D-001140-01-20

13778075
VCA-1001
H7904

9806

F10348
IP-202-1012
RS-122-2301
E7370
EGMK81312

GSE130306
GSE130306
GSE130306
GSE130306
GSE130306
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www2.replicationdomain.com

GEO GSM432687
GEO GSM429321
GEO GSE86747
GEO GSE98448
GEO GSE98448
GEO GSE98448
GEO GSE98448
ENAPRJEB8073
ENA PRJEB8073
GSE87039
GSM923444

N/A

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https://bioconductor.org/packages/release/bioc/html/DESeq2.html
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http://smithlabresearch.org/software/methpipe/
https://github.com/dekkerlab/cworld-dekker
https://www.graphpad.com/
https://www.multiscalegenomics.eu/MuGVRE/
http://smithlabresearch.org/software/methpipe/
Supplementary Figure 3

A. DO/cycling

B. Percentage of LADs vs. distance of SAHD to the membrane (µm)

C. SAHD compartments

D. TR scenario

E. Frequency vs. distance of SAHD to the membrane (µm)

F. SAHD compartments

G. MR scenario

H. Frequency vs. distance of SAHD to the membrane (µm)

I. SAHD compartments

J. MR- WI Scenario

K. Frequency vs. distance of SAHD to the membrane (µm)

L. SAHD compartments

M. Density vs. genomic distance (bp)

N. Spearman=0.66

O. Distance (TADbit) (µm) vs. Distance (polymer) (µm)
Supplementary Figure 6

A. Comparison of Average Insulation Score (AIS) in OISD4 cells with siNT and siDNMT1.

B. Log2(obs/exp) for OIS-D0 and OIS-D4.

C. Linear view of genome with SAHF organisation.

D. Venn diagram showing genes upregulated in OIS4 cells and genes in SAHD adjacent regions.

E. FPKM expression levels for CXCR4, GPR39, RHOB, and OTOF in OIS-D0 and OIS-D6.

F. p-value ranges for molecular and cellular functions.

G. Diagram showing chromosome 6 with SAHD, SAHF, PNLDC1, IGF2R regions.

H. Immunofluorescence staining for SAHF, IGF2R, and PNLDC1.

I. Interprobe distance (µm) for IGF2R - PNLDC1, PNLDC1 - SAHF, IGF2R - SAHF.

J. FPKM expression levels for IGF2R and PNLDC1 in OIS-D0 and OIS-D6.