Near-atomic resolution structures of interdigitated nucleosome fibres

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Chromosome structure at the multi-nucleosomal level has remained ambiguous in spite of its central role in epigenetic regulation and genome dynamics. Recent investigations of chromatin architecture portray diverse modes of interaction within and between nucleosome chains, but how this is realized at the atomic level is unclear. Here we present near-atomic resolution crystal structures of nucleosome fibres that assemble from cohesive-ended dinucleosomes with and without linker histone. As opposed to adopting folded helical ‘30 nm’ structures, the fibres instead assume open zigzag conformations that are interdigitated with one another. Zigzag conformations obviate extreme bending of the linker DNA, while linker DNA size (nucleosome repeat length) dictates fibre configuration and thus fibre–fibre packing, which is supported by variable linker histone binding. This suggests that nucleosome chains have a predisposition to interdigitate with specific characteristics under condensing conditions, which rationalizes observations of local chromosome architecture and the general heterogeneity of chromatin structure.
Eukaryotic DNA is packaged by basic proteins into chromatin, which consists of repeating nucleosome units composed of a core element of ~146 base pairs (bp) wrapped around an octamer of four different core histone proteins in $\frac{1}{2}$ left-handed turns. The nucleosome core elements are connected by a variable length of typically 10–90 bp of linker DNA, yielding the nucleosome, which can be associated with a 5th type of histone protein, linker histone. The structural and chemical disposition of chromatin components beyond the DNA sequence, epigenetic features, influence DNA accessibility and nuclear factor activity, which enable control of genomic activities in the cell with temporal and spatial precision.

Condensing of nucleosome chains into compact states is modulated by several distinct types of architectural factors, including linker histones, which are essential proteins in higher eukaryotes that serve critical functions in gene regulation, maintaining genomic integrity and generating mitotic chromosomes for cell division. We have an extensive understanding of how epigenetic features of individual nucleosomes—including nucleosome structure, stability, dynamics and positioning as well as DNA methylation, core histone post-translational modifications and variant localization—influence genome activities. However, our comprehension of how nucleosomes are organized into higher order structures is comparatively lacking, and yet this is essential for understanding genomic function. The reason why our grasp of chromatin higher order organization is relatively poor compared to the structure and function of individual nucleosomes is twofold in that in vitro characterization involves studying very large assemblies that are intrinsically dynamic in nature. Second, at the cellular level, imaging techniques that afford the capacity to resolve individual nucleosomes in situ are just coming of age.

Up until the last decade or so, the textbook consensus on chromatin structure was that the subsequent compaction level of the nucleosome fibre took the form of folded, helical, so-called 30 nm, conformations, for which there are two general models, either a solenoidal or a two-start zigzag configuration. Indeed, 30 nm fibres form in vitro under carefully controlled conditions at low divalent metal cation concentrations with regularly spaced nucleosome arrays and with chromatin isolated from various cell types. However, in recent years, a number of studies applying a variety of imaging and analytical methods to in situ interphase and mitotic chromatin as well as that isolated from cells have provided evidence of compact chromatin consisting of irregular interdigitated fibres with zigzag characteristics, but not 30 nm structures. Moreover, under condensing Mg$^{2+}$ concentrations, nucleosome arrays have been shown to form large globular structures that consist of extended nucleosome fibres, which are interdigitated with one another. Nevertheless, 30 nm fibre conformations have been reported to occur in the context of terminal differentiation, consistent with special structural features associated with transcriptionally silenced chromatin.

The key questions remain as to what factors determine whether chromatin assumes interdigitated or 30 nm configurations and if the former coincides with a common mode of organization in vivo, then how is this architecture realized at the atomic level.

As a solid-state method, pioneering crystallographic investigations have yielded detailed insight into chromatin compaction and higher order structure. However, the approach is challenged by the dynamic nature of nucleosomal systems, limiting the ability to obtain well-ordered crystals of large assemblies. Indeed, few crystal structures of nucleosomes (i.e., >147 bp) have been obtained, and those reported are based on blunt-ended DNA fragments that coincide with a lack of double helix continuity in the lattice and typically significant disorder that limits resolution. Here, we present findings from cohesive-ended dinucleosomes that self-assemble into lattices composed of fibres with uninterrupted Watson–Crick continuity of the double helix. The interdigitated nucleosome fibre structures shed light on chromosome architecture and help rationalize seemingly conflicting reports on chromatin structure.

### Results

**Dinucleosomes assemble into open zigzag fibres.** We obtained nucleosome fibre crystals with two different dinucleosome constructs of 349 and 355 bp that differ with respect to the lengths of the linker DNA (Figs. 1 and 2; Supplementary Fig. 1; Supplementary Movies 1 and 2). The 145 bp nucleosome core elements of both constructs consist of a near maximal histone octamer-affinity, Widom-based sequence flanked by histone octamer-refractory poly-A/T elements in the linker DNA arms, followed by mixed sequence elements that terminate in single-stranded TGCA overhangs. In divalent metal-containing buffers, the dinucleosomes assemble into a lattice in which the four-nucleotide cohesive termini associate through Watson–Crick base pairing (Figs. 1 and 2; Supplementary Fig. 2). The offset pairing of dinucleosomes in an open-ended fashion yields uninterrupted continuity of the DNA double helix from one end of the crystal to the other, making the lattice like a fabric woven of chromatin fibres that are interdigitated with one another. The length of the fibres composing the lattice are estimated to be up to roughly 10$^5$ nucleosomes or 18 million bp.

Nucleosome fibre structures for the 349 and 355 constructs were solved at up to 3.4 Å and 3.8 Å resolution, respectively, and in both cases the asymmetric unit is a single dinucleosome repeat (Figs. 1 and 2; Tables 1–3; Supplementary Fig. 1; Supplementary Movies 1 and 2). Fibres of the 355 construct are highly extended, having an average rise of 42.5 Å per nucleosome, with the dinucleosome repeats situated perpendicular to the fibre axis and stacked edgewise one on top of the other to create a pseudo-twofold symmetric double-sided staircase. The 31 bp ‘shared’ linker DNA sections that connect the two nucleosomes within a dinucleosome cross over roughly orthogonal to the fibre axis, whereas the 34 bp ‘paired’ (annealed) linker DNA sections that connect adjacent dinucleosomes are aligned more coincident with the fibre axis. As such, the fibre has a decisive zigzag configuration comprising two parallel nucleosome columns, where each nucleosome connects to two others in the opposing column. Due to the extended nature of the fibre, the stacking interactions between nucleosomes in a given column are sparse, but entail both divalent metal assisted DNA-DNA interdigitation and histone-DNA contacts involving symmetry-related elements at the nucleosome core periphery (Fig. 3).

In spite of differing by 2 and 4 fewer bp, respectively, in the shared and paired linker DNA sections, the 349 nucleosome fibre is analogous to that of the 355 in that it has a zigzag configuration comprising two parallel nucleosome columns (Figs. 1 and 2). However, the two fibre types are otherwise remarkably distinct. Instead of the fibre axis running between the two nucleosomes of the dinucleosome repeats as occurs for 355, the 349 fibre axis runs between the repeats, which stack with one another in an offset fashion within each column. In this regard, the nearly planar 349 dinucleosomes are cantled along the fibre axis at a roughly 45° angle, supporting an alternating arrangement where each dinucleosome is paired in turn with two different dinucleosomes of the opposing column. Moreover, one face of each nucleosome is stacked against the face of a neighbouring nucleosome with which it is connected DNA-wise via an intervening dinucleosome from the other column. In this way, the 349 fibre is in essence somewhat folded and is thus less extended than that of the 355,
with an average rise of 30.3 Å per nucleosome. In fact, the orientation of the 30 bp paired linker DNA segments is such that they run substantially retrograde (i.e., negative rise) to the overall translational sense of the fibre. The intrafibre nucleosome face-to-face contacts of the 349 configuration involve histone-histone and histone-DNA contacts as well as divalent metal-mediated interactions (Fig. 3).

Linker DNA dictates fibre structure. Given that the nucleosome core regions have identical DNA sequences and secondary structures, the pronounced differences between the 349 and 355 fibre configurations illustrate how the linker DNA dictates fibre arrangement. The influence of the linker DNA structure can be understood in two separate respects—DNA length/twist relationship as opposed to bending of the double helix. Since the two constructs have identical DNA sequences with the exception of the, relative to 349, introduction of 2/4 additional bp of mixed sequence to the shared/paired linker arms of 355 (Fig. 1), differences in bending are a result of context dependency, as opposed to sequence-specific effects. On the other hand, the bp insertions of 355 coincide with double helix twist extensions, relative to the 349 fibre, of 54° (shared) and 88° (paired) between the respectively linked nucleosome core regions. As such, the angular disposition between linked nucleosomes is highly distinct for the two fibre structures (Figs. 1 and 2; Supplementary Movies 1 and 2). Nucleosomes connected by the shared linker arms are nearly co-planar in 349 as opposed to being oblique in 355. The ~90° difference in the paired linker arms means that, while the 349 shared linker DNA sections criss-cross one another in very close proximity (<8 Å; separated by 2 intervening nucleosomes) at right angles, those of the 355 are widely separated and run parallel to one another. Therefore, the twist differences imposed by differing linker DNA lengths alone have a dominating effect on fibre configuration.

In addition to dramatic rotational distinctions brought about by the small changes in linker DNA length between the 349 and 355 fibres, there are also significant differences in bending of the linker DNA (Figs. 1 and 2). However, as outlined above, these structural differences appear to result from the interplay of orientational constraints imposed by linker DNA length and the optimization of nucleosome packing interactions. In this regard, linker DNA bending deformations are more pronounced in the 355 compared to the 349 fibre, and in fact each linker section has 5–7 flexible (pyrimidine-purine) bp steps between the flanking poly-A/T tracts. In the 349 fibre, the shared linker section displays a gentle arcing, with a 43° bend between the entry/exit points of two linked nucleosome core regions, while the paired linker section is nearly straight (19° bend). The corresponding shared and paired linker arms in the 355 fibre show an overall bending of 26° and 60°, respectively, but both arms also display some excess curvature as the double helix wriggles. By allowing interfibre groove-groove interdigititation, this translational writhing in the 355 linker DNA facilitates close packing of the fibres (see below). Moreover, the pronounced curvature in the 355 paired linker arm may help accommodate the greater difference between its shared and paired linkers (31 versus 34 bp) compared to those in the 349 (29 versus 30 bp).

Fibres integrate and interdigitate. In both the 349 and 355 systems, the fibres making up the lattice run parallel to and are
interdigitated with one another (Fig. 4; Supplementary Movies 3 and 4). Several features are common to both systems, including offset overlapping of the fibres, such that only one nucleosome column of a given fibre stacks against that of a neighbouring fibre. The other is that the columns of both fibre types display a staircase-like profile that permits a key in lock style of fibre–fibre integration. As such, the similar packing configurations between fibres comprise nucleosomal face-to-edge contacts whereby the interacting nucleosomes have roughly perpendicular orientations with respect to one another (Fig. 5a). These inter-fibre interfaces involve histone-DNA contacts as well as divalent metal-mediated interactions (Fig. 5b).

Aside from the interdigitation features common to both the 349 and 355 fibres, one notable distinction relates to the inter-fibre register of the nucleosome columns (Fig. 4; Supplementary Movies 3 and 4). For the 349, the nucleosome cores are juxtaposed with one another such that the linker DNA sections comprise an isolated zone between the two nucleosome columns within a fibre. On the other hand, the 355 nucleosome cores are offset with one another, so that there are contacts between the nucleosome cores of one fibre and the linker DNA of neighbouring fibres. As mentioned above, this coincides with distinct DNA bending conformations assumed by the 355 linker arms.

**Fig. 2 Structures of nucleosome fibres.** 12-nucleosome sections of the 349 and 355 fibres are shown. The DNA is coloured to distinguish the dinucleosome repeats. The cores histones are shown in pale yellow (H2A), pink (H2B), purple (H3) and green (H4). The inset depicts the Watson–Crick base pairing from annealing of the cohesive ends between dinucleosomes, generating the paired linker DNA sections.

**Linker histone mediates intra- and inter-fibre contacts.** We grew crystals of 349 and 355 assembled without linker histone and with saturating amounts of the H1.0 linker histone variant (Tables 1–3; Supplementary Figs. 3 and 4). In either case, fibre structure and packing are nearly identical (r.m.s.d. of superimposed dinucleosomes assembled with and without linker histone is 0.62 and 0.24 Å for the 349 and 355 systems, respectively), suggesting that a dominant role is played by the nucleosome packing and nucleosome-nucleosome interaction forces. For the 355 system, linker histone binding coincides with either dynamic and/or static (mixed binding modes) disorder, such that there is insufficient clarity of the electron density to model the linker histones. In the 349 crystals, we observe two distinct linker histone binding modes, one of which corresponds to the on-dyad mode observed in several earlier structures of nucleosome-linker histone assemblies31,32,34 (Fig. 6; Supplementary Movie 5). For this binding configuration, the linker histone globular domain sits at the nucleosome centre (dyad), where it interacts extensively
with the outward facing minor groove in addition to making contacts with both linker DNA arms. Given the close proximity of the shared linker DNA arm in the opposing nucleosome column of the fibre, the linker histone globular domain in fact makes additional contacts with this element of the juxtaposed dinucleosome. As such, the on-dyad linker histone mediates inter-nucleosomal, albeit infrafibre, interactions.

In contrast to the on-dyad association, the other linker histone binding site is remote from any nucleosome dyad region and does not involve interactions with linker DNA (Fig. 6; Supplementary Movies 5 and 6). In this ‘non-dyad’ mode, the linker histone globular domain instead sits in a niche surrounded by nucleosomes, where it interacts with nucleosome core DNA elements from three separate fibres. In this regard, the non-dyad binding linker histone mediates interfibre contacts and thereby helps consolidate the fabric of the interdigitation network.

In comparing the two different linker histone binding modes observed in the 349 fibres, it is notable that in both cases very similar DNA binding motifs are utilized by H1.0 (Fig. 6a, b). This is consistent with the positioning of the double helix elements provided by the nucleosomes from the three fibres in the non-dyad mode mimicking closely the configuration of the two linker DNA plus dyad double helix components coinciding with on-dyad binding. In fact, the nucleosome core regions from the fibres that surround the linker histone in the non-dyad mode could enhance the interaction potential by providing additional DNA binding surfaces.

**Discussion**

Here, we have shown that two distinct cohesive-ended dinucleosomes both self-assemble into continuous nucleosome chains under compacting conditions. The chains assume open configurations, in which the DNA path zigzags to form two nucleosome columns on either side of the fibre axis, permitting dense packing of nucleosomes by virtue of the staircase-like fibre profiles that foster a key in lock style of interdigitation. The structures afford a near-atomic resolution picture of how chromatin fibres can adopt

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**Table 1 Data collection and refinement statistics for 349 dinucleosome assembled with linker histone (H1.0) and crystals stabilized in either a low (L) or high concentration (H) of cryoprotectant.**

|                | 349-H1.0-L | 349-H1.0-H |
|----------------|------------|------------|
| **Data collection** |            |            |
| Space group     | P2;2;2;1   | P2;2;2;1   |
| Cell dimensions |             |            |
| a, b, c (Å)     | 121.06, 184.07, 223.37 | 120.78, 172.88, 216.04 |
| α, β, γ (°)     | 90, 90, 90 | 90, 90, 90 |
| Resolution (Å)  | 3.40–49.32 (3.40–3.59) | 3.70–90.01 (3.70–3.90) |
| Rmerge (%)      | 15.1 (248) | 21.2 (164) |
| Rfree (%)       | 4.4 (72.7) | 7.0 (57.5) |
| CC (%)          | 121 (1.1)  | 7.8 (1.4)  |
| Completeness (%)| 99.8 (98.6) | 99.7 (94.2) |
| Redundancy (%)  | 13.3 (13.1) | 10.6 (9.0) |
| **Refinement**  |            |            |
| Resolution (Å)  | 3.40–49.32 | 3.70–91.78 |
| No. reflections | 67,687     | 46,780     |
| Rwork/Rfree (%) | 20.6/26.5  | 19.4/26.6  |
| No. atoms       | 27,058     | 25,760     |
| Core histone    | 12,149     | 12,075     |
| Linker histone  | 561        | 1,128      |
| DNA            | 14,311     | 14,311     |
| Solvent         | 30         | 46         |
| B-factors (Å²) | 30         | 46         |
| R.m.s. deviations |            |            |
| Bond lengths (Å) | 0.006      | 0.006      |
| Bond angles (°) | 1.26       | 1.44       |

*Single crystal data sets; Values in parentheses are for the highest resolution shell.

**Table 2 Data collection and refinement statistics for 355 dinucleosome crystals stabilized in either a low (L) or high concentration (H) of cryoprotectant.**

|                  | 355-L | 355-H |
|------------------|-------|-------|
| **Data collection** |       |       |
| Space group      | P2_1  | P2_2  |
| Cell dimensions |       |       |
| a, b, c (Å)     | 88.99, 228.97, 118.84 | 85.04, 218.60, 115.93 |
| α, β, γ (°)     | 90, 92.68, 90 | 90, 92.32, 90 |
| Resolution (Å)  | 4.6–49.28 (4.6–4.92) | 3.81–47.98 (3.81–3.96) |
| Rmerge (%)      | 12.9 (196) | 9.6 (145) |
| Rfree (%)       | 5.4 (81.8) | 4.0 (64.6) |
| I/iR             | 6.6 (1.2) | 11.4 (1.2) |
| CC (%)          | 99.5 (48.1) | 99.9 (42.4) |
| Completeness (%)| 99.9 (100) | 99.8 (98.7) |
| Redundancy (%)  | 6.9 (6.6) | 6.8 (5.8) |
| **Refinement**  |       |       |
| Resolution (Å)  | 4.6–49.28 | 3.81–47.98 |
| No. reflections | 25,862 | 40,456 |
| Rwork/Rfree (%) | 20.5/26.1 | 20.8/28.6 |
| No. atoms       | 26,471 | 26,496 |
| Core histone    | 11,911 | 11,934 |
| Linker histone  | –      | –      |
| DNA            | 14,557 | 14,557 |
| Solvent         | 3      | 3      |
| B-factors (Å²) | 269     | 202    |
| R.m.s. deviations |       |       |
| Bond lengths (Å) | 0.005  | 0.004  |
| Bond angles (°) | 1.28   | 1.27   |

*Single crystal data sets; Values in parentheses are for the highest resolution shell.

**Table 3 Data collection and refinement statistics for 349 dinucleosome and 355 dinucleosome assembled with linker histone (H1.0), with crystals stabilized in a low (L) concentration of cryoprotectant.**

|                  | 349-L | 355-H1.0-L |
|------------------|-------|------------|
| **Data collection** |       |            |
| Space group      | P2;2;2;1 | P2_1       |
| Cell dimensions |       |            |
| a, b, c (Å)     | 120.81, 182.83, 219.03 | 89.13, 229.61, 119.05 |
| α, β, γ (°)     | 90, 90, 90 | 90, 92.71, 90 |
| Resolution (Å)  | 5.10–49.11 (5.10–5.59) | 4.90–229.6 (4.90–5.29) |
| Rmerge (%)      | 25.3 (149) | 9.4 (184) |
| Rfree (%)       | 10.5 (61.7) | 4.0 (83.0) |
| CC (%)          | 6.3 (1.4) | 9.3 (1.0) |
| Completeness (%)| 99.7 (100) | 99.5 (97.7) |
| Redundancy (%)  | 6.7 (6.7) | 6.6 (5.7) |
| **Refinement**  |       |            |
| Resolution (Å)  | 5.10–49.11 | 4.90–118.9 |
| No. reflections | 19,927 | 21,435 |
| Rwork/Rfree (%) | 22.2/28.6 | 19.5/24.6 |
| No. atoms       | 26,460 | 26,468 |
| Core histone    | 12,149 | 11,911 |
| Linker histone  | –      | –      |
| DNA            | 14,311 | 14,311 |
| Solvent         | –      | –      |
| B-factors (Å²) | 196    | 291     |
| R.m.s. deviations |       |       |
| Bond lengths (Å) | 0.005  | 0.005   |
| Bond angles (°) | 1.39   | 1.27    |

*Single crystal data sets; Values in parentheses are for the highest resolution shell.*
zigzag configurations that interdigitate with one another to produce highly condensed states. This is consistent in an overall sense with investigations of heterochromatin and metaphase chromosome architecture, which show chromatin chain interdigitation5,15–18,23 with zigzag characteristics of the individual fibres19,21,22,24. Furthermore, the chromosome-like, globular nucleosome array condensates that form under compacting divalent metal cation concentrations, similar to the ionic conditions used to obtain and stabilize the present fibre crystals, are also composed of interdigitated chains that would appear to be zigzag7,10. From the structures here, the global preference for zigzag fibre conformations is explained by obviating, or at least minimizing, the need for extreme bending of the linker DNA, as such arrangements allow for the nucleosome core entry/exit points to roughly face one another.

While the two fibre structures, 349 and 355, have similar overall features in terms of their zigzag nature and mode of interdigitation, at the local level they are highly distinct. This is a consequence of differences in the lengths of the linker DNA that connect the nucleosome core regions. Although these differences correspond to only several bp per linker arm, by altering the twist of the double helix at the radian level, they have a profound influence in particular on the orientational relationships between nucleosomes within a fibre and consequently between fibres as well. In this respect, there is an obvious interplay between the fibre configuration adopted and optimization of fibre–fibre interactions (packing), especially given their open, unfolded (355) or only partially folded (349) conformations.

By extrapolation from what we have observed here with two different nucleosomal constructs, there may be many fibre configurations associated with distinct linker DNA lengths. Linker DNA length varies depending on species, cell type, cell status and genomic location, displaying a non-uniform distribution between about 10 and 90 bp3,39. This corresponds to a broad distribution of nucleosome size, or nucleosome repeat length, ranging over roughly 160–240 bp. With consideration of only variations in linker DNA length alone, this would suggest on average a prodigious variability in local chain conformation from one section of nucleosomes to the next. We envision that nucleosome chains composed of mixed linker DNA lengths would interdigitate with each other through maximization of stacking interactions by exploiting the inherent flexibility of open (non-folded) fibres. With further consideration of variations in nucleosome core composition and the presence of other nuclear factors, this could

Fig. 3 Intrafibre nucleosome-nucleosome interactions. Nucleosome-nucleosome interfaces within the 349 and 355 fibres are shown with dashed lines indicating hydrogen bonds (black) and coordinate bonds (magenta; Ca2+ ions, grey spheres). The cores histones are shown in pale yellow (H2A), pink (H2B), purple (H3) and green (H4).
account for the observed heterogeneity of chromatin structure at the multi-nucleosomal level\textsuperscript{5,20}. Indeed, we find contacts of all sorts that stabilize nucleosome-nucleosome interactions in the fibre structures, which include the core histones, linker histones, DNA and divalent metal ions. While the highly basic N-terminal tail domains of the core histones are for the most part too disordered to model in the fibres, they have been seen to mediate a variety of internucleosomal contacts in crystal structures of nucleosome core particles\textsuperscript{40}. Therefore, the histone tails are likely to mediate potentially numerous interactions with neighbouring nucleosomes within and between fibres, as we observe for instance with H4 R23 in the 349 structure (Fig. 5b). Moreover, non-tail core histone elements, in particular the protruding C-terminal α-helix extension of H2B, stabilize specific intra- and inter-fibre interactions (Figs. 3 and 5b), which could be impacted by post-translational modifications (e.g., H2B K120 acetylation/ubiquitination). As such, in vivo, variations in histone variant composition, post-translational modifications and the presence of architectural factors, in addition to the activity of other cationic species, would be expected to have a significant impact on fibre structure or dynamics in a localized fashion, as has been suggested by a multitude of chromatin investigations.

Our results support the premise that chromatin chains have a predisposition to interdigitate by adopting zigzag conformations, and this is consistent with the present general consensus on cellular nucleosome organization\textsuperscript{3,7}. Furthermore, with the system studied here, we find that this tendency of nucleosome chains to interdigitate is unaltered by the presence or absence of linker histone, which is observed to support interdigitation configurations through binding in variable modes. Although it remains to be established what linker histone binding modes are prevalent in vivo, the findings here suggest that these architectural factors could bind opportunistically in structural niches to stabilize compact chromatin. Nonetheless, there are conflicting reports in the literature regarding whether or not chromatin can assume folded helical 30 nm structures in vivo. Some experts have suggested that 30 nm fibre structures may exist only transiently or in the context of specialized regulatory features. This would be consistent with reports of 30 nm structures coinciding with states of terminal differentiation\textsuperscript{25–28}. Moreover, our findings here

**Fig. 4 Interdigitation of nucleosome fibres.** 12-nucleosome sections for each of three integrated fibres are shown from two perspectives for the 349 and 355 systems. The central fibre is coloured magenta, with the two adjacent fibres in yellow and cyan.
emphasize the critical role played by the length of the linker DNA, and consequently its regularity from nucleosome to nucleosome. Studies on 30 nm structures have shown that folding is also dependent on linker DNA size\textsuperscript{9,41,42}, and this is consistent with reports of 30 nm fibre structures present in centromeric regions where nucleosomes are positioned in well phased regular arrays\textsuperscript{43}. Alternatively, the presence or action of other nuclear factors—not just histone and architectural proteins but possibly also chromatin remodelling activities—may be involved in promoting 30 nm fibre folding.

In recent times, there have been a number of breakthrough investigations shedding light on nucleosome organization in vivo. Some of these works utilize direct imaging methods while others rely on sophisticated proximity mapping strategies that yield internucleosomal contact relationships on the order of 100s of bp in separation\textsuperscript{19,24,29}. The 349 and 355 fibre models are in remarkable agreement with these proximity profiles in showing enrichment of N to N\textsuperscript{+2} contacts (interactions between nucleosomes N and N + 2; 349 and 355) but also of N to N + 3 contacts (349 only), albeit to a lesser extent (Fig. 7). Importantly,
the fibre models fit the 'α-structure' (tetrahedron) and 'β-structure' (rhombus) chromatin folding motifs determined for yeast and provide a means to fit/interpret a proximity map of human cellular chromatin without the need for invoking folded helical 30 nm structures. Nevertheless, the human proximity map revealed exciting distinctions in structure between open (non-compact), H3K9me3-marked silenced and H3K27me3-marked silenced chromatin. In fact, the 355 fibre structure is remarkably similar to the 2-start zigzag form of 30 nm structure. Both share dominant N + 2 contact relationships from the zigzag arrangement of the chain, but the 355 configuration (246 Å height, 140 Å width and 552 Å length over 12 nucleosomes) is as if one had
stretched out (and untwisted) the 2-start 30 nm fibre (272 Å height, 272 Å width and 287 Å length over 12 nucleosomes) so as to largely unfold it. In summary, the structures and approach presented here may help to provide a more unified understanding of local chromatin organization and dynamics as well as offer a means for further atomic-level characterization of nucleosome compaction and architectural factor activities.

Methods
Production of DNA fragments. The non-palindromic 349 and 355 DNA fragments consist respectively of 345 bp and 351 bp duplexes with 4-nucleotide overhangs at each 3′ terminus (Fig. 1). Three tandem repeats of the 349 and 355 DNA fragments were cloned into the pUC19 vector with KpnI sites flanking each repeat (GenScript, Piscataway, NJ, USA). Subsequent to bacterial overexpression and plasmid preparation, the DNA fragments were cleaved from the vector by digestion with KpnI. This was followed by FPLC purification using a Resource Q column (GE Healthcare, Chicago, IL, USA) to remove vector. The sequences of both DNA strands are as follows:

349-Strand 1
CGCTGGAAAAAAAAAACGCATCCCGGTGCCGAGGCCGCTCAATTGGTCGTAGACAGCTCAGTGAGCGTCTCAACGCGTCC
CTTAACCGCAGCTACGGCAGGCTCTCACAGCGGTGACCTAAGCGCTACGCGGTCTGCAGCTCTACGCGCCGCACATGAAAAAA
ACCAGCGGTAC

349-Strand 2
CGCTGGTTTTTTTTTTCATGTGCCGGTCTCACACGTGCCTGGAGACTAGTAAGCGCTTCTAGTGGCGGTTAAAACGCGGTAGACAGCGCGTACGTGCGTTTAAGCGGTGCTAGAGCTGTCTACGACCAATTGAGCGGCCTCGGCCACGGGCACATGAAAAAATGACACGCGGTAC

355-Strand 1
CGCTGACGAAAAAAAAAACGCATCCCGGTGCCGAGGCCGCTCAATTGGTCGTAGACAGCTCAGTGAGCGTCTCAACGCGTCC
CTTAACCGCAGCTACGGCAGGCTCTCACAGCGGTGACCTAAGCGCTACGCGGTCTGCAGCTCTACGCGCCGCACATGAAAAAA
ACCAGCGGTAC

355-Strand 2
CGCTGACGTTTTTTTTTTCATGTGCCGGTCTCACACGTGCCTGGAGACTAGTAAGCGCTTCTAGTGGCGGTTAAAACGCGGTAGACAGCGCGTACGTGCGTTTAAGCGGTGCTAGAGCTGTCTACGACCAATTGAGCGGCCTCGGCCACGGGCACATGAAAAAATGACACGCGGTAC

Fig. 7 Agreement between nucleosome fibre structures and nucleosomal proximity mapping studies of chromatin. Four-nucleosome sections of the 349 and 355 fibres are shown on the left, with the double helix coloured by dinucleosome repeat (as shown in Fig. 2). Nucleosome-nucleosome proximity relationships (primarily N to N + 2, secondarily N to N + 3) found to be enriched in chromatin mapping studies19,24,29 are indicated. On the right, for comparison, is a schematic representation of the folding motifs found in yeast chromatin24.
Linker histone production. *Homo sapiens* H1.0 was cloned into the pET11b vector with an N-terminal hexa-histidine tag (EZBiolab Inc., Carmel, IN, USA). Linkers were expressed in BL21(DE3) or BL21(DE3)amiY (Escherichia coli) cells cultured in 2×TY medium at 37 °C. Subsequent to the cell density reaching an OD$_{600}$ of 0.6, protein expression was induced by adding IPTG (isopropyl β-D-1-thiogalactopyranoside) to a final concentration of 0.5 mM and allowed to proceed for 3–4 h at 37 °C or overnight at 18 °C. Cells were harvested by centrifugation, and the pellets resuspended in lysis buffer (50 mM Tris-HCl [pH 8.0], 250 mM NaCl, 0.05% [v/v] Calbiochem protease inhibitor cocktail set III—EDTA-free [EMD Millipore, Billerica, MA, USA]) with 4 mg/ml lysozyme, followed by stirring at 4 °C for 1 h. Subsequent to centrifugation at 4 °C, the pellet was resuspended in lysis buffer containing 1 M NaCl. After sonication on ice, the solution was allowed to incubate with stirring at 4 °C overnight. Subsequent to centrifugation at 4 °C, the supernatant, which contains linker histone, was retained.

Linker histone was purified starting with FPLC affinity-tag purification using a HiTrap HP column (GE Healthcare, Chicago, IL, USA). Heavy DNA contamination was removed by washing the protein-loaded HisTrap column with HisTrap HP column (GE Healthcare, Chicago, IL, USA). Supernatant, which contains linker histone, was retained. Incubate with stirring at 4 °C overnight. Subsequent to centrifugation at 4 °C, the supernatant, which contains linker histone, was retained. H. sapiens H1.0 was expressed in BL21(DE3) or BL21(DE3)amiY (Escherichia coli) cells cultured in 2×TY medium at 37 °C. Subsequent to the cell density reaching an OD$_{600}$ of 0.6, protein expression was induced by adding IPTG (isopropyl β-D-1-thiogalactopyranoside) to a final concentration of 0.5 mM and allowed to proceed for 3–4 h at 37 °C or overnight at 18 °C. Cells were harvested by centrifugation, and the pellets resuspended in lysis buffer (50 mM Tris-HCl [pH 8.0], 250 mM NaCl, 0.05% [v/v] Calbiochem protease inhibitor cocktail set III—EDTA-free [EMD Millipore, Billerica, MA, USA]) with 4 mg/ml lysozyme, followed by stirring at 4 °C for 1 h. Subsequent to centrifugation at 4 °C, the pellet was resuspended in lysis buffer containing 1 M NaCl. After sonication on ice, the solution was allowed to incubate with stirring at 4 °C overnight. Subsequent to centrifugation at 4 °C, the supernatant, which contains linker histone, was retained.

**Dinucleosome and linker histone complex assembly.** Recombinational constructs and linker histone complexes were assembled with recombinant *H. sapiens* core histones $^{44,45}$ and either the 349 or 355 DNA fragment. The core histones with N-terminal hexa-histidine tags were expressed in BL21(DE3) (H2A, H2B and H3) or JM109(DE3) (H4) *E. coli* cells and affinity purified with a HiTrap IMAC FF column (GE Healthcare, Chicago, IL, USA). Linker histone-rich fractions were subsequently pooled and the N-terminal His-tag was removed by the addition of human rhinovirus 3C protease (Thermo Fisher Scientific, Waltham, MA, USA). The His-tag cleaved linker histone sample was subjected to size-exclusion chromatography on a HiLoad 16/60 Superdex 200 column (GE Healthcare, Chicago, IL, USA). Linker histone complexes were reconstituted based on established protocols$^{46}$ and shown to be capable of assembling into long fibre structures in solution (Supplementary Fig. S1). Incubation of linker histones, linker histone 349 and 355, was mixed with nuclease in nuclease staining solution (Supplementary Fig. 6).

**Dinucleosome ligation assay.** In total, 349 and 355 dinucleosome (2 μM) samples were ligated using 40 Units of T4 ligase (NEB, Ipswich, Massachusetts, USA) in ligation buffer containing 50 mM Tris (pH 7.5), 10 mM MgCl$_2$, 1 mM adenosine-5′-diphosphate, and 1 μM n-tRNA$_{HIV}$ (Ambion, Austin, TX, USA) in a 4 μl reaction volume. The reaction was incubated at room temperature, and for each time point (0.5, 1, 2 or 3 h) 10 μl sample were withdrawn. In total, 349 and 355 dinucleosome (2 μM) samples were ligated using 40 Units of T4 ligase (NEB, Ipswich, Massachusetts, USA) in ligation buffer containing 50 mM Tris (pH 7.5), 10 mM MgCl$_2$, 1 mM adenosine-5′-diphosphate, and 1 μM n-tRNA$_{HIV}$ (Ambion, Austin, TX, USA) in a 4 μl reaction volume. The reaction was incubated at room temperature, and for each time point (0.5, 1, 2 or 3 h) 10 μl sample were withdrawn. The reaction was stopped with 25 mM EDTA followed by subjection to 0.8% agarose gel electrophoresis (using a running buffer [0.25X TBE] of 22.3 mM Tris, 22.5 mM boric acid and 0.5 mM EDTA). Gels were stained with ethidium bromide for DNA visualization. The procedure described above was repeated with four independent experiments; a representative result is shown in Supplementary Fig. 5.

**Crystalization and data collection.** Assemblies were prepared for crystallization by mixing linker histone and dinucleosome with a slight excess of linker histone (1.2–1.4 linker histone:nucleosome molar stoichiometry) in a buffer consisting of 80–100 μM CaCl$_2$, 50 mM KCl and 20 mM Na-acetate [pH 4.5] for 349 and 40–80 μM CaCl$_2$, 50 mM KCl and 10 mM Na-acetate [pH 4.5] for 349 and 20–40 μM CaCl$_2$, 25 mM KCl and 10 mM Na-acetate [pH 4.5] for 355, with incubation at 18 °C.

Subsequent to centrifugation at 18 °C, linker histone complexes were harvested and stabilized in a buffer consisting of 20–25 μM CaCl$_2$, 25 mM KCl and 10 mM Na-acetate [pH 4.5], with 0.2 M Na$_2$CO$_3$, 3% glycerol (v/v), 0.1 mg/ml lysozyme, and 0.05% β-mercaptoethanol (pH 7.5). Crystals of linker histone assemblies were grown by the hanging-drop vapour diffusion method through saltin-g via equilibration against a reservoir solution containing 40–50 mM CaCl$_2$, 25 mM KCl and 10 mM Na-acetate [pH 4.5] for 349 and 20–40 μM CaCl$_2$, 25 mM KCl and 10 mM Na-acetate [pH 4.5] for 355, with incubation at 18 °C. Crystals were transferred into 0.1 M CaCl$_2$, 25 mM KCl and 10 mM Na-acetate [pH 4.5] for 349 and 20–40 μM CaCl$_2$, 25 mM KCl and 10 mM Na-acetate [pH 4.5] for 355, with incubation at 18 °C.

**References**

1. McCarty, R. K. & Tan, S. Nucleosome structure and function. *Chem. Rev.* 115, 2255–2273 (2015).
2. Zhou, K., Gaulier, G. & Luger, K. Nucleosome structure and dynamics are coming of age. *Nat. Struct. Mol. Biol.* 26, 3–13 (2019).
3. Baldi, S., Korber, P. & Becker, P. B. Beads on a string—nucleosome array arrangements and folding of the chromatin fiber. *Nat. Struct. Mol. Biol.* 27, 109–118 (2020).
4. Fedorov, D. V., Zhou, B. R., Skoultchi, A. I. & Bai, Y. Emerging roles of linker histones in regulating chromatin structure and function. *Nat. Rev. Mol. Cell Biol.* 19, 192–206 (2018).
5. Ou, H. D. et al. ChromEMT: Visualizing 3D chromatin structure and compaction in interphase and mitotic cells. *Science* 357, eaag0025 (2017).
6. Luger, K., Dechassa, M. L. & Tremeirenc, D. J. New insights into nucleosome and chromatin structure: an ordered state or a disordered affair? *Nat. Rev. Mol. Cell Biol.* 13, 436–442 (2012).
7. Hansen, J. C. et al. The 10-nm chromatin fiber and its relationship to interphase chromosome organization. *Biochem. Soc. Trans.* 46, 67–76 (2018).
8. Hansen, J. C. Conformational dynamics of the chromatin fiber in solution: determinants, mechanisms, and functions. *Annu. Rev. Biophys. Biomol. Struct.* 31, 361–392 (2002).
9. Robinson, P. J., Fairall, L., Huynh, V. A. & Rhodes, D. EM measurements define the dimensions of the “30-nm” chromatin fiber: evidence for a compact, interdigitated structure. *Proc. Natl Acad. Sci. USA* 103, 6506–6511 (2006).
10. Maeshima, K. et al. Nucleosomal arrays self-assemble into supramolecular granules lacking 30-nm fibers. *EMBO J.* 35, 1115–1132 (2016).
27. Kizilyaprak, C., Spehner, D., Devys, D. & Schultz, P. In vivo chromatin organization: evidence for a complex hierarchical organization. J. Cell Biol. 105, 77–92 (1987).

28. Scheffer, M. P., Eltsov, M. & Frangakis, A. S. Evidence for short-range helical nucleosome folding motifs. Proc. Natl Acad. Sci. USA 105, 19732–19737 (2008).

29. Bednar, J. et al. Nucleosomes, linker DNA, and linker histone form a unique structural motif that directs the higher-order folding and compaction of chromatin. Proc. Natl Acad. Sci. USA 95, 14173–14178 (1998).

30. Eltsov, M., Macellan, K. M., Maashima, K., Frangakis, A. S. & Dubochet, J. Analysis of cryo-electron microscopy images does not support the existence of 30-nm chromatin fibers in mitotic chromosomes in situ. Proc. Natl Acad. Sci. USA 105, 19732–19737 (2008).

31. Fussner, E. et al. Open and closed domains in the mouse genome are configured as 10-nm chromatin fibers. EMBO Rep. 13, 992–996 (2012).

32. Bednar, J. et al. Mapping nucleosome resolution chromosome folding yeast by Micro-C. Cell 162, 108–119 (2015).

33. Ricci, M. A., Manzo, C., Garcia-Parajo, M. F., Lakadamyali, M. & Cosma, M. P. Chromatin fibers are formed by heterogeneous groups of nucleosomes in vivo. Cell 160, 1145–1158 (2015).

34. Grigoryev, S. A. et al. Hierarchical looping of zigzag nucleosome chains establishes the 30-nm chromatin structure. Proc. Natl Acad. Sci. USA 113, 1234–1243 (2016).

35. Cai, S., Bock, D., Pilhofer, M. & Gan, L. The in situ structures of mono-, di-, and trinucleosomes in human heterochromatin. Mol. Biol. Cell 29, 2450–2457 (2018).

36. Chican, A. et al. Frozen-hydrated chromatin from metaphase chromosomes has an interdigitated multilayer structure. EMBO J. 38, e9796 (2019).

37. Ohno, M. et al. Sub-nucleosomal genome structure reveals distinct nucleosome folding motifs. Cell 176, 520–534 (2019).

38. Langmore, J. P. & Schutt, C. The higher order structure of chicken erythrocyte chromatin. J. Mol. Biol. 119, 1699–1702 (1979).

39. Woodcock, C. I. Chromatin fibers observed in situ in frozen hydrated onion sections. Native fiber diameter is not correlated with nucleosome repeat length. J. Cell Biol. 125, 11–19 (1994).

40. Kizilyaprak, C., Spehner, D., Devys, D. & Schultz, P. In vivo chromatin organization of mouse rod photoreceptors correlates with histone modifications. PLoS ONE 5, e110390 (2010).

41. Scheffer, M. P., Eltsov, M. & Frangakis, A. S. Evidence for short-range helical order in the 30-nm chromatin fibers of erythrocyte nuclei. Proc. Natl Acad. Sci. USA 108, 16992–16997 (2011).

42. Risca, V. I., Denny, S. K., Straight, A. F. & Greenleaf, W. J. Variable chromatin structure revealed by in situ spatially correlated DNA cleavage mapping. Nature 541, 237–241 (2017).

43. Schalch, T., Duda, S., Sargent, D. F. & Richardson, T. J. X-ray structure of a tetranucleosome and its implications for the chromatin fiber. Nature 436, 138–141 (2005).

44. Zhou, B. R. et al. Structural mechanisms of nucleosome recognition by linker histones. Mol. Cell 59, 628–638 (2015).

45. Bednar, J. et al. Structure and dynamics of a 197 bp nucleosome in complex with linker histone H1. Mol. Cell 66, 384–397 (2017).

46. Ekundayo, B., Richardson, T. J. & Schalch, T. Capturing structural heterogeneity in chromatin fibers. J. Mol. Biol. 429, 3031–3042 (2017).

47. Zhou, B. R. et al. Revisit of reconstituted 30-nm nucleosome arrays reveals an ensemble of dynamic structures. J. Mol. Biol. 430, 3093–3110 (2018).

48. Garcia-Saez, I. et al. Structure of an H1-Bound 6-nucleosome array reveals an untwisted two-start chromatin fiber conformation. Mol. Cell 72, 902–915 (2018).

49. Lowry, P. T. & Widom, J. New DNA sequence rules for high affinity binding to histone octamer and sequence-directed nucleosome positioning. J. Mol. Biol. 276, 19–42 (1998).

50. Thäström, A., Bingham, L. M. & Widom, J. Nucleosomal locations of dominant DNA sequence motifs for histone-DNA interactions and nucleosome positioning. J. Mol. Biol. 338, 695–709 (2004).

51. Chua, E. Y., Vasudevan, D., Davey, G. E., Wu, B. & Davey, C. A. The mechanics behind DNA sequence-dependent properties of the nucleosome. Nucleic Acids Res. 40, 6358–6352 (2012).

52. Widom, J. A relationship between the helical twist of DNA and the ordered positioning of nucleosomes in all eukaryotic cells. Proc. Natl Acad. Sci. USA 89, 1095–1099 (1992).
