Evidence for the Existence of Globular Units in the Supranucleosomal Organization of Chromatin


One of the significant steps in the past decade of chromatin research has been the finding of the nucleosome as the common principle form of DNA-histone organization (Hewish and Burgoyne 1973; Kornberg 1974; Olins and Olins 1974; for review see Chambon 1978). In most eukaryotic nuclei the bulk of the chromatin is associated with histones in a regular pattern and is organized, in the non-transcribed chromatin regions, in nucleosomal particles, which can be visualized by electron microscopy of spread chromatin in extended nucleosomal filaments, i.e. 10–12 nm large beads-on-a-string (Fig. 1 a). This principle of packing in globular repeating units results in an overall foreshortening of the DNA by a factor of 5–6. However, some other forms of DNA arrangement in chromatin have also been reported: (i) the dinoflagellate chromosome (for references see Rae and Steele 1978); (ii) DNA newly replicated in the absence of protein synthesis (Riley and Weintraub 1979); (iii) actively transcribed chromatin in which the nucleohistone is organized in such a way that the contained DNA is in a predominantly extended configuration (Fig. 1 b; Foe et al. 1976; Foe 1978; Franke et al. 1976, 1978; Reeder et al. 1978; Trendelenburg et al. 1976; for references see Franke et al. 1979); and (iv) chromatin present in cells infected with adenovirus (Kedinger et al. 1978; Matsuguchi et al. 1979; Miller and Hodge 1975) and herpes simplex virus (Müller et al. 1980): In the case of herpes-infected cells, a total of four different forms of chromatin are observed (Fig. 1 c).

However, the reports of all these different types of nucleosomal and non-nucleosomal chromatin configurations are almost exclusively based on studies of chromatin dispersed in solutions containing very low salt concentrations. It is obvious that extended nucleosomal filaments as shown in Fig. 1 a do not occur in the living cell nucleus since the ion concentrations determined in nuclei of living cells (Century et al. 1970; Century and Horowitz 1974; Dick and Fry 1973; Paine et al. 1975; Riemann et al. 1969) favor the arrangement of nucleosomes into higher order structures. The important question therefore is: What is the organization of the chromatin in the nucleus of the living cell?

It has been long known that transcriptionally inactive chromatin is condensed and appears, in electron micrographs of ultrathin sections through fixed cells, in the form of densely stained 20–50 nm thread-like units, often with a periodic granular substructure. Such arrays of dense chromatin are often especially conspicuous along the inner nuclear membrane (e.g., Fig. 6 a; for references see Franke and Scheer 1974; Franke et al. 1978; Zentgraf et al. 1980a, b). Discussions on the higher order organization of chromatin can start from the general consensus that transcriptionally inactive chromatin appears in fibrillar arrays of diameters 20–50 nm but the interpretation of the arrangement of the nucleosomal chain within this higher order fibril is highly controversial. In particular, there is continuing debate whether the thick fibril seen in sections is continuous and homogeneous (Davies et al. 1974; Davies and Haynes 1976; Everid et al. 1970), or is composed of tightly apposed globular subunits of the diameter of the fiber (Franke and Scheer 1974; Franke et al. 1978; Zentgraf et al.
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1975; Zentgraf et al. 1980a, b). Recently two extreme concepts on the in vivo packing of the nucleosomal filament in the higher order fibril have been proposed: On one hand, supercoiled or solenoidal arrangements of the nucleosomal chain into a continuous, uniformly thick 25 nm fiber has been suggested (Finch and Klug 1976; Thoma et al. 1979; Worcel 1980); on the other hand, the chromatin fibril has been interpreted as an array of closely packed higher order granules of different dimensions (20–50 nm) and nucleosome contents, possibly characteristic of the specific type of chromatin (for references see Table I). In the following several lines of experimental observations will be presented which support the concept that nucleosomal chains are arranged in granular particles of diameters about two or three times the size of the nucleosome.

The first indication for the existence of granular forms of higher order packing of nucleosomal chains has come from studies on the small chromatin circles (“minichromosomes”) of intracellular forms of simian virus 40 (SV40). When prepared under physiological salt conditions, the SV40 chromatin is arranged in granular particles of about 30 nm (Fig. 2a) which, upon exposure to low salt concentrations, can be unravelled into nucleosome-containing circles (Fig. 2b; Griffith 1975; Griffith and Christiansen 1978; Keller et al. 1978; Müller et al. 1978; Zentgraf et al. 1978; see there for further references). Assembly of circular DNA molecules into nucleosomes and supranucleosomal chromatin structures is not restricted to viral chromatin. In oocytes of several organisms such as the water beetle, Dytiscus marginalis, and the house cricket, Acheta domesticus, large amounts of amplified ribosomal DNA occur in rings containing one, two or more ribosomal RNA genes (Gall and Rochaix 1974; Trendelenburg 1974; Trendelenburg et al. 1976). When these circles of rRNA genes are not transcribed as, e.g., in the nucleolar chromatin of previtellogenic oocytes, they show, upon prolonged exposure to low salt buffer, the characteristic beaded string configuration indicative of nucleosomal packing (Fig. 2c; Scheer and Zentgraf 1978). In addition, such inactive chromatin circles often appear to be compacted into globular higher order structures of diameters 21–34 nm, suggesting that each higher order granule is an aggregate of 6–9 nucleosomes (Insert in Fig. 2c; Scheer and Zentgraf 1978).

The assembly of circular DNA into chains of nucleosomal particles, and subsequently, into higher order granules can be studied experimentally. When circular DNA molecules are injected into amphibian oocyte nuclei they are assembled into chromatin-like configurations showing typical nucleosomal morphology (Wyllie et al. 1978; Zentgraf et al. 1979). After brief dispersal in low salt buffer or after incubation once at maintained physiological ionic strength, it was demonstrated that the injected DNA molecules are assem-
Fig. 2. Condensed chromatin of simian virus 40 arranged in granular 30 nm large particles (a), which can be unravelled into nucleosomes upon exposure to low salt concentrations (b). Beaded aspect indicative of nucleosomal arrangement is also observed in the transcriptionally inactive rDNA chromatin circles of previtellogenic oocytes of the water beetle, Dytiscus marginalis (c), which also can form 20–35 nm large supranucleosomal granules (Insert in Fig. 2c; for details see Scheer and Zentgraf 1978). Bars 0.2 μm. a 106,000×. b 128,000×. c 86,000×. Insert in c 84,000×
Table 1

<table>
<thead>
<tr>
<th>Reference</th>
<th>Material</th>
<th>Methods</th>
<th>Conclusion</th>
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<tbody>
<tr>
<td>Franke et al. 1976</td>
<td>Chicken erythrocytes, isolated nuclei from rat liver and cultured cells</td>
<td>Brief dispersion in low salt buffers, spread preparations</td>
<td>&quot;... these larger chromatin globules (26 nm) represent further condensation and packing of the nucleosomal bead chains, characteristic for certain topological classes of inactive chromatin ...&quot;</td>
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<td>Kiryanov et al. 1976</td>
<td>Isolated rat liver nuclei and chromatin fractions</td>
<td>Ultrathin sections and spread preparations; emphasis on influence of Mg²⁺ and EDTA</td>
<td>&quot;The basic structural unit of chromatin is a 200 Å DNP fibril consisting of stacked globules of the diameter (200 Å).&quot;</td>
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<td>Renz et al. 1977</td>
<td>Chromatin from bovine lymphocyte nuclei, including micrococal nuclease-digested chromatin fractions</td>
<td>Sedimentation in sucrose gradients, spread preparations; emphasis on involvement of histone H1</td>
<td>&quot;... chromosome fibers are 200 Å in diameter and composed of discrete globular structures.&quot;</td>
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<td>Hozier et al. 1977</td>
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<td>&quot;Individual knobs (&quot;superbeads&quot;) are arranged in tandem and have an average diameter of about 200 Å.&quot;</td>
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bled not only into nucleosomes, at the first level, but also into supranucleosomal granules of about 30 nm diameter of higher order organization. A similar transformation of circular molecules into granular higher order chromatin structures is also observed after incubation with the proteins present in 100,000×g supernatants of homogenized amphibian oocyte nuclei and then centrifuged in discontinuous sucrose gradients containing 100 mM salt (Fig. 3; for details see Scheer et al. 1980). Under such conditions it was estimated that the DNA molecules are compacted in a way resulting in an overall foreshortening by a factor of 30−50.

The concept of a supranucleosomal organization of chromatin in regularly arranged granular units has also received support from experiments using rat liver chromatin after limited digestion with micrococal nuclease. In such experiments it was demonstrated that the nucleosomal chromatin is present in large granules which can be prepared in the form of monomeric (23 nm) and dimeric (29 nm) spheroidal particles, each containing a mean of eight nucleosomes (Strättling et al. 1978). This essentially confirms the results of Renz and colleagues (Hozier et al. 1977; Renz et al. 1977) who have shown that the 200 Å chromatin fiber is composed of tandemly arranged, discrete globular 20 nm structures.

However, such demonstrations of granular chromatin aggregates of nucleosomes ("superbeads"; Hozier et al. 1977; Renz et al. 1977) in relatively short oligonucleosomal fragments is open to the criticism that these structures result from artificial collapse and aggregation. Therefore, we decided to use non-fragmented chromatin of whole nuclei to examine the gradual, salt-dependent transition from native chromatin structure to extended nucleosomal chains, and vice versa. When isolated nuclei from a diversity of animal and plant cells (e.g. HeLa cells, Fig. 4a; corn root tip cells, Fig. 4b) are lysed and briefly incubated in low salt buffer, the partly dispersed chromatin appears in characteristic granules of diameters 25−35 nm, which are often closely packed and serially arranged into thick "discontinuous" fibrils (Fig. 4a, b; for details see Zentgraf et al. 1980a,
Fig. 3. Packing of circular DNA (plasmid PTP 6AA1 shown in a with cytochrome-c spreading) during incubation with proteins present in a 100,000× g supernatant fraction from nuclei of oocytes of *Pleurodeles waltlii* and subsequent purification by sucrose gradient centrifugation (for details see Scheer et al. 1980). The newly assembled plasmid chromatin appears at physiological salt concentrations in the form of large granular particles (b), which, at low salt concentrations, are unravelled into chains of nucleosome-like structures (c–e). Bars 0.5 μm (a), 0.2 μm (b–e). a 50,500×. b 61,000×. c 61,000×. d 61,000×. e 61,000×.
Fig. 4. Electron micrographs of spread preparations of chromatin from HeLa cells (a) and meristeme cells of corn, Zea mays (b) after brief dispersal in low salt buffers. The 25–30 nm large granular structures are often linearly arranged in fibrillar arrays. Upon prolonged exposure to low salt buffer, these higher order structures are unravelled into typical extended nucleosomal chains; (this is shown, for example, for the corn root tip chromatin in c). Bars 0.5 μm. a 67,000×. b 80,000. c 166,000×
Fig. 5. Chicken erythrocyte chromatin briefly lysed in low salt buffer reveals 18–30 nm higher order granules inter-connected by strand regions of nucleosomal filaments (a) as typical of intermediate stages of unravelling of native condensed chromatin into nucleosomal chains. Maximally extended nucleosomal filament chromatin as obtained after prolonged treatment with low salt buffer (b) is rapidly recondensed in vitro after dialysis against buffers containing physiological salt concentrations and shows similar chains of serially arranged supranucleosomal granules as the native chromatin (c; for details see Zentgraf et al. 1980 b). Bars 0.5 µm. a 51,000×. b 58,000×. c 42,000×
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Upon prolonged incubation in low salt buffer, the supranucleosomal granular structures are progressively lost and the material is completely unravelled into the extended nucleosomal chains characteristic of maximally dispersed chromatin (shown in Fig. 4c for the chromatin of Zea mays).

One of the classic objects of chromatin research is the chicken erythrocyte nucleus, which is characterized by the high degree of homogeneity of its inactive and highly condensed chromatin. Examination of spread preparations of minimally dispersed chick erythrocyte chromatin, such as present after short times of incubation in low salt buffer, reveal the presence of large (18–30 nm) chromatin granules interspersed between regions of extended nucleosomal filaments (Fig. 5a). The observation of large chromatin granules, of globular structures of intermediate sizes and shapes on the same nucleosomal filament, and the progressive unravelling of this chromatin into extended nucleosomal chains in low salt suggests that the rows of large (20–30 nm) granules represent the native state of higher order packing of the nucleosomal chain (Franke et al. 1976; Zentgraf et al. 1980b; see also Pruitt and Grainger 1980). When such avian erythrocyte chromatin present as extended nucleosomal filaments (Fig. 5b) is dialyzed against buffers of elevated salt concentrations (e.g., containing 100 mM NaCl) or, alternatively, nanoto millimolar concentrations of magnesium salts (Zentgraf et al. 1980b), it rapidly and almost completely recondenses into arrays of higher order particles of similar sizes as those observed in native chromatin (Fig. 5c; for details see Zentgraf et al. 1980a). When briefly lysed in low salt buffers, this chromatin displays a relatively homogeneous composition of uniformly sized (47 nm), large supranucleosomal granules separated by short strands of nucleosomes (Fig. 7a). Upon prolonged incubation in solutions of low salt concentrations, the chromatin strands, which are often arranged in loops, show an increased number of nucleosomes located between residual, large supranucleosomal granules (Fig. 7b). This and the increasing number of intermediate-sized granular structures of irregular sizes and shapes suggest a progressive unravelling of the higher order chromatin beads (estimated contents of 20–26 nucleosomes) into the extended nucleosomal chains (Fig. 7c). Whether the formation of the extremely large supranucleosomal beads in the sea urchin sperm chromatin is related to the presence of special forms of histones H2A, H2B, and H1 (Wangh et al. 1972) and/or to the unusually high nucleosomal repeat length of 240 nucleotides of DNA (Spadafo-
Fig. 6. Electron micrographs of ultrathin sections of chicken erythrocyte chromatin present in intact cells (a), dispersed in low salt buffer (b) and after recondensation in vitro into higher order form (c). Dispersed chromatin showing extended nucleosomal filaments (arrows in b, denote some nucleosomes revealing a relatively unstained core) is recondensed at elevated ion concentrations into relatively closely packed, 20–30 nm large granules, which sometimes are intimately associated with remnants of the nuclear envelope (NE), thus resembling the aspect of the outermost layer of the condensed chromatin observed in intact erythrocyte nuclei (a; for details on fixation procedures see Zentgraf et al. 1980b). Bars 0.1 µm. a 270,000×. b 270,000×. c 270,000×

ra et al. 1976) remains to be examined. Likewise the correlation of higher order chromatin structure (for reports of differences of nucleosomes per supranucleosomal bead see also Meyer and Renz 1979) to specific histone composition and nucleosomal DNA repeat length (for avian erythrocytes see Morris 1976, and Chambon 1978) is not clear at the moment.

In discussing the significance of the granular forms of higher order organization of the nucleosomal filament, it is worth emphasizing that the supranucleosomal granules described above are seen when possible preparation artifacts are minimized, for example by the use of unsheared long chromatin strands freshly prepared by brief lysis of isolated nuclei (see also Kiryanov et al. 1976) or
Fig. 7. Sea urchin sperm chromatin, briefly lysed in low salt buffer, is mostly contained in large, uniformly sized (40–50 nm) supranucleosomal granules that in some regions, especially in the periphery, are interconnected by short intercepts of extended nucleosomal filament morphology (a). Upon further incubation in low salt concentrations the number of large supranucleosomal granules is decreased and the proportion of extended nucleosomal filaments increased suggesting progressive unravelling into the extended nucleosomal chains (b). The final state as obtained after prolonged incubation at very low ionic strength is shown in c and presents a uniform appearance of the chromatin in extended nucleosomal filaments. Bars 0.5 µm. a 58,000 x, b 38,000, c 39,000 x

intact cells. Relatively short (up to about 0.5 µm) regions of solenoid-like configuration or forms of other more tightly, close-packed, 20–30 nm chromatin fibrils (including structures described by Olins and Olins 1979; Rattner and Hamkalo 1979; Worecel 1978) have been occasionally seen in our systematic studies on progressive chromatin dispersal in solutions of lowered salt concentrations (for an example see Zentgraf et al. 1980b), indicating that such configurations are intermediates of chromatin swelling rather than native structures formed in the physiological ion environment of the living cell nucleus.

The chemical principle, that gives rise to the formation of these specific supranucleosomal arrangements in chromatin are not under-
stood. While a general involvement of histone H1 and H5 in the condensation of the nucleosome filament into higher order fibrils and granules has been indicated by a number of experiments (Keller et al. 1978; Renz et al. 1977; Thoma et al. 1979; see there for further references), the specific role and location of these histones in the higher order structure is still unknown.

It is obvious that specific biochemical and topological experiments are needed for further understanding of the packing of chromatin at the supranucleosomal level.

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