We describe a cell-free system, derived from preblastoderm Drosophila embryos, for the efficient assembly of cloned DNA into chromatin. The chromatin assembly system utilizes endogenous core histones and assembly factors and yields long arrays of regularly spaced nucleosomes with a repeat length of 180 bp. The assembly system is also capable of complementary-strand DNA synthesis accompanied by rapid nucleosome formation when the starting template is single-stranded circular DNA. Chromatin assembled with the preblastoderm embryo extract is naturally deficient in histone H1, but exogenous H1 can be incorporated during nucleosome assembly in vitro. Regular spacing of nucleosomes with or without histone H1 is sufficient to maximally repress transcription from hsp70 and fushi tarazu gene promoters. The Drosophila assembly system should be particularly useful for nucleosome assembly during DNA synthesis and for elucidating the action of transcription factors in the context of native chromatin.

Transcription in eukaryotes is generally associated with local changes of the regular chromatin structure and composition (see reference 14 for a review). While “active chromatin” as a technical term has been in existence for many years, the exact nature and determinants of these activated regions are largely unknown (28, 43). In general, sites of transcription appear to be less condensed and thus more accessible than the tightly packed bulk chromatin. Probing the accessibility of sequences with nucleases in isolated nuclei has yielded a wealth of information about the localization of regulatory sites, such as enhancers or promoters, and their potential or actual activity.

While the correlations between transcription and accompanying chromatin features are well established, it has only recently been appreciated that structural components of chromatin can be used to regulate gene activity in a highly specific manner (16, 17, 42, 45). It has been shown that nucleosomes that are positioned in vivo with regard to the underlying DNA sequence can influence transcription and replication by modulating the accessibility of regulatory sites for the binding of protein factors in vivo. Conversely, DNA-binding proteins have been described that activate or repress transcription by influencing the position of nucleosomes in the vicinity of important regulatory sequences (10, 19, 31, 37, 46). To elucidate these phenomena at the molecular level, an in vitro reconstruction of the mutual independence of transcriptional regulators and structural components of chromatin under physiological conditions is necessary.

We are interested in general aspects of transcriptional regulation by RNA polymerase II in the context of chromatin, using the fruit fly Drosophila melanogaster as a model system. A priori, Drosophila embryos should be an excellent source of factors required for nucleosome assembly; during the initial stages of embryonic development, the fly genome is replicated and packaged into chromatin once every 9 min by using a maternal pool of histones and assembly factors (11). A cell-free system for nucleosome assembly derived from early Drosophila embryos has been reported previously (25). However, this procedure has not been adopted by workers in the field, and our attempts to create an efficient assembly extract according to the published procedure were not successful. Here we report the development of a stable and very efficient cell-free system for nucleosome assembly using Drosophila embryos. The system is also able to carry out a DNA synthesis reaction that mimics lagging-strand replication. We find that assembly of nucleosomes on hsp70 and fushi tarazu DNA templates is sufficient to maximally repress transcription in vitro, with or without the concomitant assembly of histone H1.

**MATERIALS AND METHODS**

**Extract preparation.** Drosophila embryos 0 to 100 min after egg laying were rinsed in water and allowed to settle into embryo wash buffer (0.7% NaCl, 0.05% Triton X-100) on ice to arrest further development. After four to five successive collections, the pooled harvest was dechorionated. The wash buffer was decanted and replaced with wash buffer at room temperature, and the volume was adjusted to 200 ml. After the addition of 200 ml of Chlorox bleach, the embryos were stirred vigorously for 90 s, poured back into the collection sieve, and rinsed extensively with tap water. They were then allowed to settle in 1 liter of wash buffer for about 2 min, after which the supernatant (containing the chorions) was aspirated off. Four more settings were performed: one in wash buffer, two in 0.7% NaCl, and one in extract buffer [10 mM N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid (HEPES, pH 7.6), 10 mM KCl, 1.5 mM MgCl₂, 0.5 mM ethylene glycol-bis(β-aminopropyl ether)-N,N,N',N'-tetraacetic acid (EGTA), 10% glycerol, 10 mM β-glycerophosphate; 1 mM dithiothreitol and 0.2 mM phenylmethylsulfonyl fluoride, added freshly] at 4°C. The embryos in extract buffer were settled in a 60-ml glass homogenizer on ice for about 15 min, and the volume of the packed embryos was estimated (−10 to 15 ml). The supernatant was aspirated, and the embryos were homogenized by six com-
plete strokes with a Teflon pestle connected to a drill press. All further manipulations were carried out at 4°C. The homogenate was supplemented with additional 5 mM MgCl₂ from a 1 M MgCl₂ stock solution and quickly mixed (final MgCl₂ concentration, 6.5 mM). When nucleosome assembly was assayed at low MgCl₂ concentrations, no MgCl₂ was added to the embryo homogenate (final MgCl₂ concentration, 1.5 mM). Nuclei were pelleted by centrifugation for 5 min at 5,000 rpm in a JA14 rotor (Beckman). The supernatant was clarified by centrifugation for 2 h at 40,000 rpm (150,000 × g) in an SW 50.1 rotor (Beckman) and collected with a syringe by puncturing the tubes just above the solid pellet, avoiding the floating layer of lipid. Aliquots (300 to 500 μl) were frozen in liquid nitrogen. Protein concentrations were determined with the Bradford assay, using bovine serum albumin as the standard; they were usually between 20 and 30 mg/ml.

At the last settling step in the extract buffer, not all embryos settle because of changes in osmolarity. In addition, variable degrees of settling will result in corresponding variable final extract protein concentrations. To standardize the yields of the extract preparation, an alternative procedure was employed. After the settling in 0.7% NaCl, the embryo suspension was poured onto a filter paper on a Büchner funnel, filtered under vacuum, and washed twice with 100 ml of extract buffer. The damp embryos were resuspended in 1 ml of cold extract buffer per g of embryos. Homogenization and centrifugation were performed as described above. When assayed by micrococcal nuclease digestion, this procedure yielded extracts which assembled described previously (6). Homogenization and centrifugation were performed as described above. When assayed by micrococcal nuclease digestion, this procedure yielded extracts which assembled described previously (6).

**Assembly reaction.** In a standard assembly reaction, 1.5 μg of plasmid DNA was incubated at 26°C with 3 mg of embryo extract protein and the following reagents (which were added from a 10× premix): 30 mM creatine phosphate, 3 mM MgCl₂, 3 mM ATP (pH 8), 0.1 μg of creatine phosphokinase (type 1; Sigma) per ml, and 1 mM dithiothreitol. The volume was adjusted to 200 μl with extract buffer. The final conductivity in the assembly reactions is equivalent to 65 mM KCl. The protein concentration for optimal assembly was determined with the Bradford assay, using bovine serum albumin as the standard; they were usually between 20 and 30 mg/ml.

**Supercoiling analysis.** At various times during the reaction, 40 μl of assembly mixture was removed and 10 μl of 5× stop mix (2.5% Sarkosyl, 100 mM EDTA) and 1 μl of DNAse-free RNase (Boehringer Mannheim) were added; the mixture was then incubated for 15 min at 37°C. Then 6.5 μl each of 2% sodium dodecyl sulfate (SDS) and 10-μg/ml proteinase K were added and again incubated for 30 min at 37°C. The DNA was precipitated with 2 volumes of ethanol after the addition of 1 μl of glycogen (10 μg) and 45 μl of 7.5 M ammonium acetate. After centrifugation, the DNA was washed with 80% ethanol, dried under vacuum, and finally dissolved in 8 μl of TE (10 mM Tris HCl [pH 7.5], 1 mM EDTA). DNA was electrophoresed on a 1.2% agarose gel with Tris-glycine buffer (lacking ethidium bromide) for 15 h at 20 V.

**Micrococcal nuclease analysis.** To a standard assembly reaction, 6 μl of 0.1 M CaCl₂ was added and quickly mixed. A 40-μl portion of the mixture was removed and treated the same as for supercoiling analysis. To the remaining reaction, 5 μl of micrococcal nuclease (Boehringer Mannheim; 50 U/μl in extract buffer) was added. After 0.5, 2, and 8 min at room temperature, 40 μl was again removed and the digestion was terminated. An RNase treatment for up to 1 h was followed by overnight proteinase K digestion. Gel electrophoresis of the micrococcal nuclease digestion products was as described in detail by Shimamura et al. (35). Micrococcal nuclease analysis of *Drosophila* embryo chromatin was performed as described by Wu et al. (50).

**DNA templates.** In the experiments presented here, the plasmid pshp70Al185 (3,443 bp), which contains hsp70 (locus 87A) gene sequences between the XhoI site at −185 and the AccI site at +300 (+1 is the transcriptional start), was used. The XhoI-AccI fragment was filled in with T4 polymerase and cloned into the *Hinc*II site of pBluescript SK M13+ (Stratagene). Single-stranded DNA was purified from phage particles after superinfection of a bacterial culture containing the plasmid with helper phage VCSM13 (Stratagene) according to standard procedures. A variant of pshp70Al185 (hsp70 minigene) was constructed by deleting an *Aat*Il fragment (+41 to +71) as described previously (6). The plasmid carrying the fushi tarazu promoter (−950 to +151) was as described previously (6).

**Replication assay.** Conditions for the replication assay were similar to those for standard nucleosome assembly reactions, except that 750 ng of single-stranded DNA instead of double-stranded plasmid DNA was used. For “uniform labeling,” 1 μl of [α-³²P]dCTP (2,000 to 3,000 Ci/mmoll (NEN) was added. For site-specific labeling, the single-stranded DNA was first annealed with a fivefold molar excess of a ³²P-end-labeled primer (hsp70 nucleotides −185 to −153) in 20 μl of 100 mM NaCl–5 mM MgCl₂. The mixture was incubated for 5 min at 75°C, 10 min at 37°C, and 5 min at 20°C; the extract and assembly components (without radiolabeled dCTP) were then added.

**Histone purifications.** Core histones as standards for gel electrophoresis were purified from the chromatin of 0- to 20-h-old embryos according to the method of Simon and Felsenfeld (36). They were identified by their migration behavior on SDS gels (44). HI was purified from nuclei of 0- to 12-h embryos by the procedure of Crosston et al. (6). The peak fraction of the phenyl-Sepharose column was concentrated four- to fivefold by vacuum dialysis in a mini-collodion bag (Schleicher & Schuell).

**Plasmid chromatin purification.** A total of 1.5 μg of single-stranded DNA or 3 μg of double-stranded plasmid DNA was assembled for 6 h under standard conditions. The synthesis of the complementary strand was followed by the introduction of 3 μl of [α-³²P]dCTP to the samples containing single-stranded DNA. Purification of chromatin was achieved by centrifugation through a 15 to 30% sucrose gradient followed by pelleting through a 30% sucrose cushion exactly as described previously (35). The chromatin-containing fractions were identified by the incorporation of radioactivity. Silver staining of the histone gels was done as described by Wray et al. (49).

**Transcription of assembled templates.** *Drosophila* embryo transcription reactions were prepared and coupled by assembly–transcription reactions were processed as described previously (6), with the following modifications. The chromatin assembly reaction was scaled down to 50 μl in proportion to the input amount of DNA template (180 ng each of the hsp70
and ftz plasmids. To 5 μl of the assembled chromatin reaction was added 2 μl of 0.5-μg/μl pUC vector DNA and 18 ng of hsp70 minigene in 9 μl of HEMG (25 mM HEPES [pH 7.6], 0.1 mM EDTA, 12.5 mM MgCl₂, 10% glycerol). Four micro liters of unshocked embryo transcription extract (6) and 5 μl of the following components (final concentrations) were added: 500 μM (each) ATP, GTP, CTP, and UTP; 4 mM creatine phosphate; 10 ng of creatine phosphokinase; 5 U of RNase; 4 mM dithiothreitol. The 25-μl reaction mixture was then incubated for 20 min at 26°C. DNA products were purified and analyzed by primer extension (6).

Quantification of the radioactivity was performed on dried polyacrylamide gel slices by liquid scintillation counting.

RESULTS

In vitro assembly of nucleosomes with regular spacing on plasmid DNA. In order to minimize the dilution of cytoplasmic components and to reduce the extraction of nuclear proteins, the procedure for preparing the Drosophila nucleosome assembly extract involves the homogenization of staged, preblastoderm embryos in a small volume of low-salt extraction buffer. Nuclei are pelleted by centrifugation, and the cytosolic supernatant is clarified by centrifugation at 150,000 × g (S-150). This crude S-150 extract contains the major components necessary to assemble nucleosomes on microgram quantities of plasmid DNA. When supplemented with ATP, MgCl₂, and an energy-regenerating system (3, 35), the assembly of regularly spaced nucleosomes is achieved.

The ability of the Drosophila extract to assemble nucleosomes was initially assayed by DNA supercoiling analysis (13, 23, 29). The winding of DNA around a nucleosomal core introduces about one positive superhelical turn in the DNA (13), which is rapidly relaxed by topoisomerase I activity, which is abundant in cell extracts. Deproteinization of the somatic assembly extract was initially assayed (13), which is rapidly relaxed by topoisomerase I activity, which is abundant in cell extracts. Deproteinization of the extract and DNA supercoiling was performed with Drosophila embryo extracts in the presence of increasing concentrations of KCl. Upon incubation with the S-150 extract (10 min), the negatively supercoiled plasmid was initially relaxed by topoisomerase I activity. Further incubation resulted in the reintroduction of DNA supercoils, suggesting the assembly of nucleosomal or nucleosomelike structures. DNA supercoiling was found to be inhibited by the presence of KCl concentrations greater than that already present in a standard reaction (the standard salt concentration is equivalent in conductivity to 65 mM KCl). Since the supercoiling assay was performed with a crude extract, it is difficult to ascribe the inhibitory effect of increased KCl to a specific component of the reaction. The effect of KCl concentrations lower than 65 mM was not analyzed, as the assay was constrained by the salt contribution of the extract.

To assess whether the supercoiling of the plasmid was due to the assembly of nucleosomes, we subjected the assembled plasmid to digestion with micrococcal nuclease. Micrococcal nuclease cleaves chromatin in the linker DNA between the nucleosomes (27). A limit digest of chromatin typically creates DNA fragments of the size protected by a nucleosome core (146 nucleotides), while partial digests result in a ladder of fragments corresponding to oligonucleosome-sized DNAs. As shown in Fig. 2A, digestion of the assembled chromatin resulted in extremely well-resolved mono- and oligonucleosome-sized DNA fragments. A densitometer tracing revealed 19 distinct bands (Fig. 2B), suggesting the assembly of a maximum of 19 nucleosomes on some of the plasmid templates. On these templates, the average nucleosomal repeat length was calculated to be 181 bp (3,443 bp/19). Measurement of the sizes of oligonucleosomal fragments also indicated the spacing of nucleosomes at ~180-bp intervals. We have made over 12 extract preparations by this procedure with similar results and hence have adopted this as the standard protocol. The resolution of the oligonucleosomal DNA bands over the background smear was less apparent when the levels of MgCl₂ were decreased below 6.5 mM (Fig. 2C). For reasons which are unclear, extracts that were prepared by adjusting the embryo homogenate to 6.5 mM MgCl₂ prior to (Fig. 2A) rather than after (Fig. 2C) the S-150 centrifugation were superior in their potential to create extended nucleosomal arrays.

DNA synthesis and nucleosome assembly. In vivo nucleosome assembly occurs naturally on newly replicated DNA. To test for the presence of similar activities in our extracts, we substituted the plasmid DNA with single-stranded circular DNA and added [α-32P]dCTP as a precursor for DNA synthesis. Upon incubation with the extract, the complementary strand was synthesized and the bulk of the labeled material was recovered in the form of supercoiled DNA (Fig. 3A). The assembly extract apparently contains DNA primase, polymerase (α), accessory factors, RNase H, and DNA ligase activities necessary to perform lagging-strand DNA synthesis.
FIG. 2. Micrococcal nuclease analysis of reconstituted chromatin. (A) Chromatin assembled for 6 h under standard conditions was digested with micrococcal nuclease for the indicated times. The final MgCl₂ concentration during the assembly reaction was 7 mM. Purified DNA fragments were resolved on 1.5% agarose gels and stained with ethidium bromide as described by Shimamura et al. (35). Lane M shows size markers consisting of 123-bp multimers (Bethesda Research Laboratories). (B) Densitometer tracing of the 0.5-min micrococcal nuclease digest shown in panel A. The arrow indicates the direction of electrophoresis. (C) Assembly reactions containing the indicated concentrations of MgCl₂ were incubated at 26°C for 6 h. The reconstituted chromatin was digested with micrococcal nuclease for the indicated times. The positions in the gel of nicked (nc), relaxed closed (rel), linear (lin), and supercoiled (sc) plasmids are marked by arrows.

DNA synthesis. The level of DNA synthesis was found to be stimulated by the presence of MgCl₂ up to 6.5 mM. To confirm that the supercoils in the newly synthesized DNA originate from the assembly of nucleosomes, micrococcal nuclease digestion assays were performed. First, DNA synthesis was traced by the incorporation of radiolabeled dCTP as described above, in a reaction that relied on endogenous primers in the assembly extract. This uniform labeling (Fig. 3B, right panel) displayed the overall chromatin assembly on the plasmid. Alternatively, a specific 32P-end-labeled primer was annealed to the single-stranded template before the addition of the extract. In this reaction, the labeled dCTP was omitted, and the micrococcal nuclease analysis therefore reflects chromatin assembly around the site where the primer annealed (Fig. 3B, left panel). In both reactions with single-stranded templates, assembly of regularly spaced nucleosomes over a significant fraction of the templates was already evident by 30 min of assembly, as gauged by the emergence of the ladder of oligonucleosome-sized DNA fragments upon micrococcal nuclease cleavage. An increase in the duration of chromatin assembly (60 and 120 min) further enhanced the ladder of DNA fragments, indicating complete or near complete deposition of nucleosomes on the DNA template.

Histone content of in vitro assembled nucleosomes. The regular spacing of the micrococcal nuclease cleavage sites in reconstituted chromatin is indicative of the assembly of nucleosomes. To extend this finding, we determined the histone composition of the reconstituted nucleosomes. Fully assembled plasmid chromatin (with single- or double-stranded DNA as the starting material) was purified by sucrose gradient centrifugation (35). The plasmid chromatin proteins were analyzed by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and silver staining (Fig. 4). From the reactions that contained either single-stranded (lane 1) or double-stranded (lane 2) DNA, a full complement of core histones of roughly equal stoichiometry could be recovered, while a reaction that was processed in parallel but lacked DNA (lane 3) displayed cosedimenting nonhistone proteins that were not associated with chromatin. It should also be noted that histone H1 was absent from the reconstituted chromatin, as no band of the expected size (~35 kDa; see below) was observed. Preblastoderm Drosophila embryos have been previously reported to be deficient in histone H1 (9). We have also observed an apparent histone variant in reconstituted chromatin that migrates above the histone H2B band on the gel. This variant is absent or poorly represented among the late embryo histones; the late embryo histones include another apparent, minor histone variant which migrates below the histone H2A position.

---
incorporation of exogenous H1 into reconstituted chromatin. Histone H1 has been suggested to play an important role in organizing chromatin and, in particular, to contribute to the inactive state of a gene (8, 24, 34, 46, 51). Since chromatin reconstituted with the early embryo extract was deficient in H1, we have determined whether exogenous purified H1 could be incorporated into the chromatin.

H1 was purified from 0- to 12-h Drosophila embryos to near homogeneity (Fig. 5A, lane 2) and was introduced into the assembly extract prior to the addition of the DNA. As shown in Fig. 5A, incorporation of exogenous H1 is evident by SDS-PAGE analysis of the histone content of assembled chromatin purified by sucrose gradient centrifugation. Histone H1 cosedimented with the reconstituted chromatin when the assembly was carried out with double-stranded plasmid DNA (Fig. 5A, lane 4) or single-stranded DNA (lane 3), but not in the absence of DNA (lane 5).

Furthermore, when chromatin reconstituted in the presence of H1 was analyzed by micrococcal nuclease digestion, an increased nucleosome repeat length was observed (Fig. 5B). A densitometer tracing (Fig. 5C, upper panel) showed that the nucleosome repeat length was increased from 180 bp in the absence of H1 to ~197 bp when H1 was present; this length is essentially identical to the in vivo repeat length for postblastoderm chromatin (Fig. 5C, lower panel).

Transcriptional repression of DNA templates assembled in chromatin. In order to assess the transcriptional potential of chromatin templates assembled in vitro, two DNA templates carrying the hsp70 and fushi tarazu gene promoters were assembled in the presence or absence of exogenous histone H1 and assayed by in vitro transcription. A naked, hsp70 minigene template was also introduced into the transcription mixture as a free DNA control. As shown in Fig. 6, incorporating nucleosome assembly resulted in the progressive repression of transcription from both hsp70 and fushi tarazu templates. By 60 min of chromatin assembly, when the DNA templates were assembled in a regular nucleosomal array as gauged by the appearance of a defined ladder of DNA fragments after micrococcal nuclease digestion (data not available).

FIG. 3. (A) DNA synthesis in embryo extracts. Single-stranded, circular DNA was incubated in a standard assembly reaction in the presence of MgCl₂ (concentrations as indicated) and [α-³²P]dCTP. Aliquots of the reaction were analyzed for DNA supercoiling by electrophoresis and autoradiography after the indicated periods of incubation. (B) Micrococcal nuclease analysis of newly synthesized plasmid DNA. Single-stranded DNA (660 ng) was incubated with 2.6 mg of embryo extract in a 150-µl reaction mixture in the presence of 7 mM MgCl₂. Left: α-³²P-labeled oligonucleotide primer was annealed to the template prior to the addition of the extract. Radioactive dCTP was omitted from the reaction. Right: DNA synthesis relied on endogenous primers and was traced with radioactive dCTP. After incubation for the indicated times, the reactions were adjusted to 3 mM CaCl₂ and digested with 190 U of micrococcal nuclease (Boehringer Mannheim) for 0, 2, and 8 min (arrows indicate increasing digestion). DNA was analyzed by gel electrophoresis and autoradiography. The gel on the left was exposed 18 times longer than the gel on the right. The positions in the gel of nicked (nc), relaxed closed (rel), and linear (lin) plasmids are marked by arrows.

FIG. 4. Histone composition of reconstituted chromatin. Proteins that cosedimented with assembled plasmids on a sucrose gradient were separated by SDS-PAGE and stained with silver. Nucleosome assembly reactions contained either single-stranded DNA (lane 1), double-stranded DNA (lane 2), or no DNA (lane 3) as the starting material. The positions of protein size standards are indicated to the left. Lanes H: Marker core histones from chromatin of 0- to 20-h embryos.

H1 was purified from 0- to 12-h Drosophila embryos to near homogeneity (Fig. 5A, lane 2) and was introduced into the assembly extract prior to the addition of the DNA. As shown in Fig. 5A, incorporation of exogenous H1 is evident by SDS-PAGE analysis of the histone content of assembled chromatin purified by sucrose gradient centrifugation. Histone H1 cosedimented with the reconstituted chromatin when the assembly was carried out with double-stranded plasmid DNA (Fig. 5A, lane 4) or single-stranded DNA (lane 3), but not in the absence of DNA (lane 5).

Furthermore, when chromatin reconstituted in the presence of H1 was analyzed by micrococcal nuclease digestion, an increased nucleosome repeat length was observed (Fig. 5B). A densitometer tracing (Fig. 5C, upper panel) showed that the nucleosome repeat length was increased from 180 bp in the absence of H1 to ~197 bp when H1 was present; this length is essentially identical to the in vivo repeat length for postblastoderm chromatin (Fig. 5C, lower panel).

Transcriptional repression of DNA templates assembled in chromatin. In order to assess the transcriptional potential of chromatin templates assembled in vitro, two DNA templates carrying the hsp70 and fushi tarazu gene promoters were assembled in the presence or absence of exogenous histone H1 and assayed by in vitro transcription. A naked, hsp70 minigene template was also introduced into the transcription mixture as a free DNA control. As shown in Fig. 6, incorporating nucleosome assembly resulted in the progressive repression of transcription from both hsp70 and fushi tarazu templates. By 60 min of chromatin assembly, when the DNA templates were assembled in a regular nucleosomal array as gauged by the appearance of a defined ladder of DNA fragments after micrococcal nuclease digestion (data not available).
FIG. 5. Incorporation of exogenous histone H1 into assembled chromatin. (A) Chromatin assembly reactions (400 μl) that contained either single-stranded DNA (lane 3), double-stranded DNA (lane 4), or no DNA (lane 5) and 15 μl of H1 were fractionated by sucrose gradient centrifugation. Proteins that cosedimented with the assembled minichromosomes (or the equivalent fractions in the reaction lacking DNA) were pelleted through a sucrose cushion, analyzed by SDS-PAGE, and silver stained. Lanes 1 and 6, core histones purified from the chromatin of 0- to 20-h embryos; lane 2, histone H1 (5 μl). The staining of the histone H2A band in lanes 3 and 4 is low, compared with that of the histone H2B band. (B) Micrococcal nuclease analysis of 1.5 μg of plasmid DNA, assembled as described in the Fig. 2A legend for 6 h in the absence or presence of 12 μl of H1. Size markers are as in Fig. 2A. (C) Upper panel: densitometer tracing of the lanes displaying the 2-min digestion products of panel B. The tracings were aligned with respect to the neighboring marker lanes to illustrate the change in repeat length. Lower panel: nuclei isolated from 6- to 18-h embryos were digested with micrococcal nuclease, and the digestion products were analyzed on a 1.3% agarose gel. Densitometer tracings of two lanes showing different extents of digestion are provided.

DISCUSSION

We have characterized an in vitro nucleosome assembly extract from preblastoderm Drosophila embryos. Early Drosophila embryos replicate their genomes on average once every 9 min (11) and assemble the newly synthesized DNA into chromatin by using the maternal pool of histones and assembly factors. They are thus an excellent source of the components required for replication and chromatin assembly. The extraction procedure we have employed was adapted from recent studies of in vitro chromatin assembly that used Xenopus oocytes as the starting material (34, 35). The Drosophila assembly system is characterized by a number of features that render it a highly useful alternative for studies on chromatin assembly. (i) Microgram amounts of plasmid DNA can be quantitatively assembled into chromatin carrying the maximal number of regularly spaced nucleosomes. (ii) The nucleosomes assembled in vitro re-
As in the *Xenopus* system (1–4), the *Drosophila* extract can be used either to assemble nucleosomes onto double-stranded plasmid DNA or in conjunction with DNA synthesis starting from a single-stranded template. Chromatin that does or does not contain histone H1 can be obtained. (iv) Large quantities of extracts with high activities are easily and reproducibly prepared according to the standard protocol. In particular, we have not encountered the seasonal variation in the activity of extracts prepared from frog oocytes. The extracts can be kept for many months at −80°C and can be thawed and frozen several times without noticeable loss of activity.

There are several technical points important to the overall activity of the assembly extract that should be noted. The *Drosophila* embryos are homogenized in a low volume of extraction buffer in order to minimize the dilution of cytoplasmic assembly components and, presumably, to approach physiological assembly conditions. The extract buffer contains low salt, 10 mM KCl, which minimizes extraction of nuclear proteins into the cytosol. The extended centrifugation at 150,000 × g in the presence of 5 mM magnesium pellets ribosome subunits from the extract that would otherwise obscure the analysis of plasmid chromatin proteins (35). The low ionic strength of the extraction buffer also enables the assembly reaction to be carried out in the presence of an energy regenerating system (ATP and creatine phosphate) and 7 mM MgCl₂, which are necessary for the assembly of long arrays of spaced nucleosomes. Importantly, the assembly relies entirely on the endogenous pool of maternal histones and their carrier proteins and assembly factors, the native state and stoichiometry of which is maintained. This may be crucial for the reconstitution of long arrays of spaced nucleosomes on natural DNA sequences, a property lacking in chromatin reconstituted by using polyanions as an assembly vehicle for exogenous histones (e.g., see references 26, 33, and 41 and references therein).

The pool of components required for proper nucleosome assembly should be highest in the early embryos and should be rapidly depleted as replication proceeds. Indeed, older *Drosophila* embryos (4 to 6 h after egg laying) yielded significantly weaker extracts (data not shown). Extracts from *Xenopus* oocytes, which also contain a maternal histone pool, are similarly efficient in quantitatively assembling large amounts of plasmid DNA into chromatin while maintaining a high degree of regular nucleosomal spacing (34, 35).

Tissue culture cells may not be as rich in assembly components, which suggests why cell-free chromatin assembly systems derived from mammalian cells appear to be less potent (5, 15).

In agreement with previous reports (2, 5, 30), elevated concentrations of MgCl₂ are required to reconstitute extensive arrays of spaced nucleosomes in our cell-free system (Fig. 2). Interestingly, we have observed rapid supercoiling of plasmid DNA in the extract at protein concentrations insufficient for the generation of spaced nucleosomes and at a suboptimal MgCl₂ concentration (1.5 mM; data not shown). Whether this supercoiling is a result of the winding of the DNA around subnucleosomal particles, such as H3-H4 tetramers, remains to be established. Such particles are likely to be intermediates in the assembly of nucleosome cores (1, 12, 18, 26, 32, 39).

The lack of histone H1 in chromatin reconstituted by the early embryo extract is not due to a deficiency of the
assembly machinery, since exogenous H1 can be efficiently assembled, resulting in an increased linker length. The nucleosomal spacing of chromatin assembled in the presence of exogenous H1 is very similar to the natural spacing of bulk chromatin in postblastoderm embryos. Hence, chromatin assembled with exogenous histone H1 in vitro closely approximates the physiological structure. Although H1 assembly is a determinant of the average linker length between core particles, it appears not to be required for regular spacing of nucleosomes on the DNA itself. Such a function may be dependent on other factors present in the extract, analogous to those found in the Xenopus system and mammalian systems (32, 38). It will be of interest to further determine at nucleotide resolution whether the specific positions of nucleosomes assembled in vitro under physiological conditions are exactly equivalent to those found in vivo or to those reconstituted by using other procedures, such as salt dialysis.

A recent study with chromatin templates reconstituted with core histones and histone H1 has implied a significant role for histone H1 in the repression of transcription by RNA polymerase II (24). The present results with the hsp70 and fushi tarazu promoters, and previous reconstitution studies with Xenopus oocyte extracts, indicate, however, that the optimal assembly of regularly spaced nucleosome cores is sufficient to maximally repress transcription in vitro, even in the absence of histone H1. It is probable that the different effects observed with histone H1 are due to the different procedures employed for core histone deposition, which lead to differences in the spacing of reconstituted nucleosomes. It is also possible that a requirement for repression by histone H1 is specific to the individual promoter sequences used in the separate studies.

The ability to assemble different types of chromatin under near-replicative conditions in vitro opens avenues for studying the competition between histone deposition during DNA synthesis and the binding of factors that govern transcriptional activation. Moreover, the preblastoderm cytoplasmic extract itself does not support RNA polymerase II transcription (data not shown), in contrast to the nuclear extracts derived from later embryos which are highly active for transcription (20, 40). An extract for nucleosome reconstitution that is free of interfering transcription initiation by polymerase II should be a particularly useful alternative to the assembly systems derived from Xenopus eggs and oocytes for studies on the interrelationships between transcriptional regulators and chromatin structure (6, 7, 22, 47, 48).

ACKNOWLEDGMENTS

We thank A. Wolff for helpful suggestions and G. Wall for excellent technical assistance with the transcription experiments. We also thank two referees for helpful comments.

P.B.B. was supported by a fellowship from the Fogarty International Center.

REFERENCES


