Electron Microscopic Visualization of Sites of Nascent DNA Synthesis by Streptavidin–Gold Binding to Biotinylated Nucleotides Incorporated In Vivo

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Abstract. Biotinylated nucleotides (bio-11-dCTP, bio-11-dUTP, and bio-7-dATP) were microinjected into unfertilized and fertilized *Xenopus laevis* eggs. The amounts introduced were comparable to in vivo deoxynucleoside triphosphate pools. At various times after microinjection, DNA was extracted from eggs or embryos and subjected to electrophoresis on agarose gels. Newly synthesized biotinylated DNA was analyzed by Southern transfer and visualized using either the BluGENE or Detek-hrp streptavidin-based nucleic acid detection systems. Quantitation of the amount of biotinylated DNA observed at various times showed that the microinjected biotinylated nucleotides were efficiently incorporated in vivo, both into replicating endogenous chromosomal DNA and into replicating microinjected exogenous plasmid DNA. At least one biotinylated nucleotide could be incorporated in vivo for every eight nucleotides of DNA synthesized. Control experiments also showed that heavily biotinylated DNA was not subject to detectable DNA repair during early embryogenesis (for at least 5 h after activation of the eggs). The incorporated biotinylated nucleotides were visualized by electron microscopy by using streptavidin–colloidal gold or streptavidin–ferritin conjugates to bind specifically to the biotin groups projecting from the newly replicated DNA. The incorporated biotinylated nucleotides were thus made visible as electron-dense spots on the underlying DNA molecules. Biotinylated nucleotides separated by 20–50 bases could be resolved. We conclude that nascent DNA synthesized in vivo in *Xenopus laevis* eggs can be visualized efficiently and specifically using the techniques described.

The ability to easily localize, at high resolution, sites of nascent DNA synthesis on DNA molecules replicating in vivo would represent a major technical advance relevant to many areas of cell biology. Currently, single- and double-stranded DNA molecules can be readily distinguished from each other by electron microscopy (15). This technique does not, however, distinguish between nascent and preexisting DNA. Newly synthesized DNA can be visualized as a track of silver grains by light microscopy using DNA fiber autoradiography (24, 25; reviewed in references 16 and 23). The DNA strands that give rise to the grain tracks, however, cannot be resolved. Nascent DNA has been observed also by electron microscope DNA fiber autoradiography (38). This technique is technically demanding and has only limited resolution (better than 0.25 μM; i.e., 750 bp).

The high affinity of avidin (20) and streptavidin (14) for biotin provides a theoretical basis for visualizing nascent DNA in the electron microscope. Biotinylated analogs (29) of deoxyribonucleoside triphosphates, the normal precursors of DNA synthesis, and electron-dense streptavidin–colloidal gold and streptavidin–ferritin conjugates have recently become commercially available. Prokaryotic and eukaryotic DNA polymerases have been shown to incorporate biotinylated nucleoside triphosphates into DNA in vitro at 30–40% of the normal rate (29). Moreover, incorporated biotinylated nucleotides can be visualized on gels using commercially available streptavidin-based detection systems. These systems have been used successfully to detect specific DNA sequences with biotinylated DNA or RNA probes (30). Avidin– and streptavidin–gold conjugates also have been used for the high resolution electron microscopy of biotinylated probes hybridized in situ to specific sequences on metaphase chromosomes (21, 35).

To use biotinylated nucleotides to visualize nascent DNA synthesized in vivo, it is first necessary to show that the biotinylated derivatives do not significantly perturb normal DNA replication. Several criteria must be satisfied. (a) The biotinylated nucleotides must be incorporated in vivo efficiently with kinetics similar to the incorporation of unbiotinylated nucleotides; (b) the incorporated nucleotides must not be removed efficiently by DNA repair processes in vivo (i.e., the newly replicated DNA containing the biotinylated nucleotides must remain intact long enough to be visualized); and (c) the incorporated nucleotides must be detectable at...
high resolution by electron microscopy. We demonstrate here that all of these conditions are met in microinjection experiments using Xenopus laevis embryos.

Materials and Methods

Reagents

Biotin-11-dUTP, biotin-7-dATP, qX174 DNA, Hind III restriction endonuclease, BluGENE1 (Bethesda Research Laboratories, Gaithersburg, MD) nonradioactive nucleic acid detection system, and streptavidin–collodial gold conjugate (20 nm) were obtained from Bethesda Research Laboratories (Gaithersburg, MD). Streptavidin–collodial gold conjugates (5 and 10 nm) were from Sigma Chemical Co. (St. Louis, MO). Bio-11-CTP and Detekt-hrp kits were from Enzo Biochem Inc. (New York, NY). 32P-labeled dATP and TTP (3000 Ci/mmol) were from ICN Radiochemicals (Irvine, CA). pXgB1 (a gift from R. Patient), pX108, and pBR322 were purified in our laboratory by standard procedures (32). Oligonucleotide primers were synthesized on a DNA Synthesizer (model No. 8750; Biosearch, San Rafael, CA) and were purified by HPLC on a Vydac C4 column (I) by the Iowa State University Nucleic Acid Facility. Escherichia coli DNA polymerase I large fragment was from Promega Biotech (Madison, WI).

Microinjection of Nucleotides and DNA

Adult Xenopus laevis were acquired from Xenopus I (Ann Arbor, MI). Healthy female frogs were injected with human chorionic gonadotropin at least 12 h before they were stripped for eggs. The eggs were fertilized in 5% De Boers solution according to the procedure of Wolf and Hedrick (42). Both fertilized and unfertilized eggs were dejellied with 2% cysteine solution (pH 7.7) before they were microinjected according to the protocol of Gurdon (22). Unfertilized eggs were irradiated for 30 s with ultraviolet light (254 nm) to inactivate the female pronucleus before microinjection. Biotinylated and 32P-labeled nucleoside triphosphates were injected into the animal hemisphere with and without exogenous plasmid DNA in a total volume of 50 nl. The microinjected eggs and embryos were incubated for the specified times at 22°C in modified Barth's solution containing 5% Ficoll type 400 (Sigma Chemical Co.) (36).

Electron Microscopy and Detection of Incorporated Nucleotides Using Streptavidin–Collodial Gold or Streptavidin–Ferritin Conjugates

DNA was further purified by HPLC on a Nucleogen DE-4000-lU column or by column chromatography on benzoylated naphthoylated DEAEE cellulose (Boehringer Mannheim Biochemicals, Indianapolis, IN), or on low-temperature melting agarose gels. The purified DNA was suspended either in 10 mM Hepes (Sigma Chemical Co.) buffer (pH 7.5) containing 100 mM NaCl or in 100 mM Tris–HCl buffer (pH 7.5) containing 150 mM NaCl. Suitable concentrations of streptavidin–collodial gold or streptavidin–ferritin conjugates were determined empirically for each sample of DNA. Samples were titrated with free biotin, when required, to reduce the number of ligand binding sites on the streptavidin–collodial gold conjugates. The streptavidin–collodial gold or streptavidin–ferritin conjugates and biotinylated DNAs were allowed to react for 15 min at room temperature and then spread for electron microscopy using the microdrop procedure of Lang and Mitani (28) or with the Kleinschmidt spreading procedures (27) using cytochrome c (15). Grids were stained with uranyl acetate, rotary shadowed with Pt-Pd (80:20), and photographed on Zeiss EM 109 or Phillips 420 electron microscope, usually at a magnification of 7,000-20,000.

Results

Biotinylated Nucleotides Were Incorporated Efficiently into Plasmid and Endogenous High Molecular Weight DNA In Vivo

As shown in Fig. 1 a, biotinylated nucleotides were incorporated into both endogenous high molecular weight (chromosomal) DNA and plasmid DNA microinjected into unfertilized eggs and embryos. The rate of incorporation of biotinylated nucleotides closely paralleled the incorporation of coincjected 32P-TTP in the same eggs and embryos (compare Fig. 1 a with the corresponding autoradiogram shown in Fig. 1 b). Embryos microinjected with up to 10 pmol each of bio-11-dCTP and bio-11-dUTP developed normally through at least the midblastula stage, although some exhibited abnormal morphology during late gastrulation. Incorporation into endogenous high molecular weight (chromosomal) DNA, which was present at far lower concentrations (6 pg/nucleus vs. 4 ng plasmid DNA/embryo), was nonetheless much more efficient than incorporation into microinjected plasmid DNA with either biotinylated or 32P-labeled nucleoside triphosphates (see Discussion).

Microinjected single-stranded circular qX174 (+) strand DNA also served as an excellent template in unfertilized eggs for incorporation of biotinylated nucleoside triphosphates (Fig. 2). Within 1.5 h substantial incorporation of biotinylated nucleotides into molecules comigrating with fully double-stranded Form II DNA was observed. By 3 h after microinjection, DNA comigrating with Form I (supercoiled) qX174 DNA was the most heavily biotinylated species. These kinetics were not significantly different from those of single-stranded DNA replicated in unfertilized eggs into which biotinylated nucleoside triphosphates had not been injected (not shown).

These results establish that biotinylated nucleotides were incorporated in vivo into endogenous or plasmid DNA molecules without visibly interfering with early embryogenesis.
Figure 1. Incorporation of biotinylated nucleotides into pBR322 plasmid and endogenous high molecular weight DNA in vivo. (a) 4 ng pBR322 Form I DNA, 0.05 μCi [32P]TTP, 10 pmol bio-11-dCTP, and 10 pmol bio-11-dUTP were coinjected into either unfertilized eggs or embryos at the 1-2-cell stage. DNA was isolated after 4 h from the unfertilized eggs and after 2.75, 6.0, 8.0, and 9.0 h from the embryos. DNA was isolated and subjected to electrophoresis on 1% agarose gels as described in Materials and Methods. Each lane was loaded with DNA from 5 eggs. After electrophoresis, the DNA was transferred to nitrocellulose filters using the Southern procedure. The incorporated biotinylated nucleotides were detected using the Detek-hrp streptavidin-horseradish peroxidase nucleic acid detection system. Lane 1 contains pBR322 marker DNA, which is not visualized by the detection system. Staining conditions were chosen such that the Form I, Form II, and high molecular weight bands were clearly defined at later stages of embryogenesis. As a result synthesis at the 16-cell stage was not prominently stained. Using more sensitive detection conditions, synthesis at the 16-cell stage can be emphasized, but lanes 5 and 6 become overstained, obscuring the bands (not shown). (b) An autoradiogram of the same gel shown in a. Synthesis at the 16-cell stage is more visible here than in a.

and without grossly modifying the apparently normal replication of either the endogenous or the microinjected DNA molecules.

Biotinylated Nucleotides were Incorporated in Proportion to Their Representation in the Endogenous Deoxynucleoside Triphosphate Pools

The internal deoxynucleoside triphosphate pools in Xenopus laevis eggs have been determined by Woodland and Pestell (43) to be 13, 16, 11, and 9 pmol each for dATP, dCTP, dGTP, and TTP, respectively. We microinjected biotinylated nucleoside triphosphates in amounts ranging from 3% (Fig. 3, lane 1) to slightly in excess of 100% (Fig. 3, lane 5) of the endogenous pools. Incorporation increased in proportion to the amounts injected up to at least 3 pmol of each nucleoside triphosphate analog per egg. As greater amounts of biotinylated nucleoside triphosphates were microinjected, the mobility of the biotinylated DNA molecules in the gel decreased (Fig. 3, lane 5 especially), presumably because the increased bulk of the newly synthesized DNA retarded its migration (see Fig. 9 and Discussion). It is important to emphasize that synthesis of φX174 DNA appeared to be normal even when the amounts of biotinylated nucleoside triphosphates microinjected exceeded the endogenous dNTP pools. We did note in other experiments, however, that survival of embryos microinjected with amounts of biotinylated nucleo-
Quantitation of the Efficiency of Incorporation of Biotinylated Nucleotides In Vivo

To quantitate the efficiency of incorporation of biotinylated nucleotides, we compared the amount of biotinylated nucleotides incorporated in a defined in vitro reaction from which the unbiotinylated homologue had been omitted vs. the amount incorporated using the same template in vivo. Synthesis in vitro was carried out using primed single-stranded φX174(+) strand DNA and the Klenow fragment of E. coli DNA polymerase I in a reaction from which TTP was omitted and replaced entirely by bio-11-dUTP (see Materials and Methods). The newly synthesized strands of the fully double-stranded Form II DNA molecules, which migrated as a discrete band on an agarose gel, were ~24% biotinylated (i.e., all dTMPs were replaced by bio-dUMPs).

The amount of incorporated biotinylated nucleotides detected using the BluGENE (Bethesda Research Laboratories) kit on known (ethidium bromide–stained bands compared to a titration of DNA of known concentration) amounts of DNA synthesized in vitro was then visually compared to the amount detected on DNA synthesized in microinjected eggs in vivo (Fig. 4). 15 pmol each of bio-7-dATP, bio-11-dCTP, and bio-11-dUTP were microinjected in the in vivo experiment.

To quantitate the efficiency of incorporation we assumed that both strands of the Form II DNA synthesized in vivo were biotinylated. This could lead to an underestimate of the efficiency of biotinylation for those molecules in which only one strand had been synthesized in vivo. Thus, our calculation of the efficiency of incorporation could be low by a factor approaching two. We also assumed that all three biotinylated nucleoside triphosphates were incorporated at similar efficiencies (i.e., that bio-11-dCTP substituted for dCTP and bio-7-dATP for dATP with at least the same efficiency that bio-11-dUTP substituted for TTP).

Fortuitously, comparable amounts of Form II DNA (monitored by ethidium bromide staining) synthesized in vivo and in vitro exhibited comparable staining for biotinylated nucleotides (legend to Fig. 4). We therefore calculated the efficiency of incorporation of biotinylated nucleotides in vivo to be at least one-sixth of the unbiotinylated (endogenous) nucleotide incorporation rate. This rate is based on the presumption that the DNA synthesized in vitro contained only one strand synthesized with only one biotinylated nucleoside triphosphate precursor, whereas the DNA syn-
Figure 4. Quantitation of the efficiency of incorporation of biotinylated nucleotides in vivo. The amount of biotinylated nucleotide incorporated during conversion of single-stranded bacteriophage ϕX174 DNA to double-stranded ϕX174 Form II DNA was compared to the amount incorporated into Form II DNA in vivo. Lanes A–J contain dilutions of the reaction product synthesized in vitro. Lane K is an unbiotinylated ϕX174 marker. Lanes L–T contain dilutions of the product synthesized in vivo using the reaction conditions in Fig. 3, lane 5. The amount of Form II DNA in each lane was quantitated by comparing the intensity of the ethidium bromide-stained DNA band to that of a band of known concentration run in an adjacent lane. The amounts of Form II added in lanes A–T are as follows: 30, 8, 3, 1.5, 0.3, 0.15, 0.03, 0.015, 0.007, 0.0035, 200 (standard), 0.06, 0.012, 0.025, 0.05, 0.25, 0.5, 1.25, 2.5, and 5 ng, respectively. Incorporated biotinylated nucleotides were visualized with the BluGENE detection system. (Only relative amounts of incorporated biotinylated nucleotides in each Form II band were compared visually. Form I DNA is not made in vitro and therefore cannot be compared to Form I made in vivo.) For example, the Form II band in lane G (0.03 ng synthesized in vitro) is less intensely stained than the band in lane N (0.025 ng synthesized in vivo). Since the DNA in lane G was 24% biotinylated on one strand (12% if both strands are considered), the Form II DNA in lane N must have incorporated >12% biotinylated nucleotides. Similarly, the Form II band in lane C (3 ng synthesized in vitro) is as intensely stained as the Form II band in lane S (2.5 ng synthesized in vivo), again indicating that the Form II band formed in vivo must contain at least 12% biotinylated nucleotides.

Stability of Heavily Biotinylated DNA In Vivo

Heavy biotinylated DNA (every TTP on one strand replaced with a biotinylated dUTP) might be expected (see Discussion) to be subject to extensive repair processes in unfertilized eggs and embryos. To test this hypothesis, biotinylated ϕX174 Form II DNA synthesized in vitro (see Materials and Methods) was microinjected into unfertilized Xenopus laevis eggs along with [32P]dATP. As shown in Fig. 5 a, microinjected biotinylated Form II DNA was not degraded for at least 2 h in the eggs. Instead, the heavily biotinylated Form II DNA was efficiently converted to Form I (supercoiled) DNA and to higher molecular weight forms, as is unbiotinylated DNA (7, 8, 17, 39). Identical results were obtained in microinjected embryos (Fig. 1 and unpublished observations). As shown in Fig. 5 b for unfertilized eggs and Fig. 1 b for embryos, incorporation of [32P]dATP was into the same plasmid forms with similar kinetics. 5 h after microinjection, a small decrease in biotinylated DNA perhaps may have occurred in both the unfertilized eggs (Fig. 5 a) and embryos (not shown). This may not be a physiological repair process, since some of the microinjected unfertilized eggs and embryos also become necrotic around this time. We conclude, therefore, that even heavily biotinylated DNA is sufficiently stable to persist, undegraded, in unfertilized eggs and in developing Xenopus laevis embryos for at least several hours. The biotinylated DNA cannot, therefore, be subject to extensive DNA repair during this time period. We also conclude that the parental (biotinylated) and newly synthesized daughter (32P-labeled) strands are converted to the same forms with similar kinetics.

Electron Microscopic Visualization of Incorporated Biotinylated Nucleotides by Streptavidin–Colloidal Gold and Streptavidin–Ferritin Conjugates

To visualize the sites of incorporation of biotinylated nucleotides, DNA isolated either from in vitro reactions or from in vivo synthesis was incubated with streptavidin–colloidal gold or streptavidin–ferritin conjugates. The DNA was spread for electron microscopy using either the microdrop procedure (28) or Kleinschmidt spreading procedure as modified by Davis et al. (15). To demonstrate the specificity of incorporation, Hind III restriction fragments of pX108 were digested briefly with exonuclease III, then end-labeled by fill-in synthesis using Klenow fragment with bio-I-dUTP substituted for TTP. As shown in Fig. 6 a, the streptavidin–colloidal gold conjugate specifically labeled the ends of the restriction fragments. No colloidal gold particles were
observed at internal locations within the restriction fragment or associated with unbiotinylated plasmid DNA molecules included as a control in the spreading mixture.

Similar binding specificity to biotinylated DNA was observed when an entire molecule contained biotinylated nucleotides. Complementary strands to φX174(+)-strand DNA were synthesized using bio-11-dUTP to replace the TTP in an in vitro reaction as described in Materials and Methods. The biotinylated double-stranded φX174 DNA was mixed with unbiotinylated double-stranded pXGαB1 DNA (17 kb) then reacted with streptavidin-colloidal gold and prepared for electron microscopy. Fig. 6 b shows that the biotinylated φX174 DNA was specifically associated with streptavidin-colloidal gold, whereas little or no gold was associated with the control unbiotinylated pXGαB1 DNA. It was also clear, however, that binding of the streptavidin-colloidal gold interfered with the proper spreading of the DNA molecules. In part, these difficulties were due to the size and high density of the gold particles. In addition, all commercially available streptavidin-gold conjugates have a large number of potential biotin binding sites (for example, ~64 potential sites on a 20-nm gold particle). This apparently caused both inter- and intrastrand cross-linking of the biotinylated DNA molecules. To reduce the cross-linking, titration of the streptavidin-colloidal gold with free biotin was carried out before the binding reaction. The binding of streptavidin-colloidal gold to biotinylated φX174 DNA before and after complete titration of the biotin binding sites is shown in Fig. 7, a and b. None of the gold particles were able to bind the biotinylated DNA in the latter case. This demonstrates that the streptavidin-colloidal gold binds to biotin moieties attached to the DNA. In contrast, untilated particles caused clumping (Fig. 7 a) and extensive cross-linking of the biotinylated DNA (Fig. 7 c). We then attempted to reduce the number of potential binding sites per gold particle to an average of one. Biotinylated φX174 circular and linear DNA molecules after reacting with titrated streptavidin-colloidal gold conjugates (20-nm size) are shown in Fig. 8, a and b. Fig. 8 c shows the association of streptavidin-colloidal gold particles (20 nm) with long linear DNA molecules gap-filled with biotinylated nucleotides. A higher magnification electron micrograph of a heavily biotinylated small fragment associated with closely spaced streptavidin-colloidal gold conjugates (20-nm size) is shown in Fig. 8 d. The closest spacing between the colloidal gold particles on this fragment is ~50 bp, approaching the theoretical limit for a 20-nm particle. Since the gold particles apparently impeded spreading (i.e., long fragments labeled heavily with 20-nm gold particles, as in Fig. 8 d, do not spread well) we tested streptavidin-colloidal gold conjugates of smaller size (5 nm) as well as streptavidin-ferritin conjugates (5-7 nm). Fig. 8 e shows the association of streptavidin-ferritin conjugates with biotinylated DNA. Although the contrast of the bound streptavidin-ferritin conjugates was not as striking as with the 20-nm colloidal gold particles, they could still be distinctly visualized. Moreover, the smaller particles allowed better spreading of the DNA, especially with longer molecules. Further experiments are in progress to increase the contrast and decrease the background using streptavidin-ferritin conjugates; these conjugates have a potential resolution of 10-15 bp.

Discussion

In this paper we describe a method for visualizing nascent DNA synthesized in vivo. It is important to emphasize that
Figure 6. Electron microscopic visualization of specific binding of streptavidin–colloidal gold conjugate (20-nm size) to biotinylated DNA. (a) End-labeled restriction fragments. Hind III–digested pX108 plasmid DNA was biotinylated as described in Materials and Methods. The biotinylated DNA fragments were mixed with unbiotinylated supercoiled plasmid DNA and labeled with colloidal gold conjugate, spread by the droplet procedure, picked up on glow-discharged carbon grids, and processed for electron microscopy. Colloidal gold was found specifically associated with the ends of the restriction fragments only. (b) Uniformly labeled φX174 DNA. Biotinylated φX174 DNA was prepared as described in Materials and Methods. A mixture of biotinylated φX174 DNA (5.4 kb) along with unbiotinylated pXGαβ1 DNA (17 kb) was labeled with streptavidin–colloidal gold conjugate, spread for electron microscopy under aqueous conditions, and photographed on a Zeiss EM 109 microscope. Colloidal gold conjugate was observed exclusively associated with 5.4 kb φX174 biotinylated DNA, but the DNA is clumped and cross-linked. Bars, 0.1 μm.

this method faithfully monitors the apparently normal incorporation of nucleoside triphosphates into chromosomal DNA during early embryogenesis, and can be used to analyze physiological replication processes. The method exploits both the high affinity of streptavidin for biotin and the high electron density of gold and ferritin particles. Avidin, a 68,000 D glycoprotein from the eggwhite (20), and streptavidin, a protein secreted by Streptomyces avidinii (14), both exhibit very high affinity for biotin ($K_a = 10^{-15}$ M). This property has been used in the development of a variety of immunobiochemical and immuno-electron microscopic techniques. For example, avidin–ferritin conjugates have been used earlier to detect, by electron microscopy, discontinuities and repair sites in linear DNAs labeled in vitro with bio-11-dUTP (26, 34). The use of avidin in sensitive detection experiments, however, has some critical limitations. Since avidin is positively charged, with $pI = 10$ at pH 7, it tends to bind nonspecifically to negatively charged molecules like nucleic acids (our unpublished observations). In addition, the glycoprotein moiety of avidin tends to bind to a variety of carbohydrates and lectins, which causes high backgrounds (our unpublished observations). Substitution of streptavidin, which has $pI = 5$ at physiological pH, for avidin, seemed to solve both the nonspecific binding and high background problems.

Use of biotinylated nucleoside triphosphate analogs by DNA polymerases in DNA synthesis has been demonstrated previously in vitro (29) and in crude nuclear extracts (9, 10). In this study we show that replicating endogenous chromosomal DNA as well as replicating exogenous DNA microinjected into Xenopus laevis eggs or embryos were efficiently labeled in vivo with biotinylated analogs of the normal precursors of DNA synthesis. The rate of incorporation of these analogs into DNA was increased proportionally by increasing the concentration of the microinjected biotinylated dNTP pools inside the unfertilized or fertilized eggs. This
suggests that the analogs successfully compete with the normal nucleoside triphosphates as substrates for DNA polymerases in vivo, in spite of the projecting biotin side chains that might be expected to interfere with incorporation (Fig. 9). Moreover, the kinetics of incorporation exactly paralleled the incorporation of unbiotinylated radioactive nucleotides in the same eggs and embryos (Fig. 1 b). The biotin-containing side chains significantly affect the bulk of the DNA molecule; this is suggested by the decreased mobility observed for heavily biotinylated DNA on agarose gels (Fig. 3, lane 5). This alteration apparently does not significantly perturb normal DNA replication.

The amount of biotinylation was proportional to the amounts of the biotinylated nucleoside triphosphates injected (Fig. 3). Incorporation of the biotinylated nucleoside triphosphates could be as high as 12% (1 in 8) of the total nucleotides incorporated. Both the endogenous chromosomal DNA and the microinjected exogenous DNA underwent multiple rounds of DNA replication. The biotinylated plasmid DNA accumulated linearly over time (based on densitometric scans of the bands in Fig. 1 a and in similar experiments), whereas the biotinylated high molecular weight DNA accumulated exponentially. This accumulation paralleled the accumulation of radioactive dNTPs in the same embryos. Moreover, the embryos underwent normal divisions at least until the late blastula or early gastrula stages, and apparently developed normally.

The incorporation levels of the analogs relative to normal substrates was somewhat lower in vivo than the 30-40% efficiency reported in vitro (29). Moreover, incorporation into endogenous high molecular weight DNA was much more efficient than into microinjected plasmid DNA (Fig. 1 a). The reason for this greater efficiency is not entirely clear. It is not related to the incorporation of biotinylated precursors of DNA synthesis, since Marini et al. (33) have similarly shown in incorporation experiments that high molecular weight DNA is replicated much more efficiently than circular plasmid DNA microinjected into the embryos (also see Fig. 1 b). The greater efficiency of replication of high molecular weight DNA does not simply result from the fact that chromosomal DNA is found in the nucleus. Forbes et al. (18), Lohka and Masui (31), and Newport (37) have shown that microinjected plasmid DNA is rapidly assembled into structures that are morphologically and functionally indis-
tistinguishable from normal nuclei. Moreover, formation of a nuclear envelope around the microinjected DNA was essential for efficient replication to occur (37).

To see whether the biotinylated nucleotides incorporated into the DNA were excised or repaired we microinjected heavily biotinylated circular DNA. Repair might have been expected a priori based on the structure of biotinylated DNA, since the biotin-containing side chain (11 atom linker) projects ~21-22 Å out of the major groove of the helix (Fig. 9). These biotins are readily accessible to binding of the electron-dense ligands, and are thus presumably also “visible” to repair enzymes. Heavily biotinylated DNA was nonetheless stable for at least the first 4–5 h after microinjection into either unfertilized eggs or embryos. This suggests that there are no repair mechanisms that quickly eliminate or repair the biotinylated nucleotides during early embryogenesis. We do not rule out the possibility that repair mechanisms work slowly or, alternatively, may be lacking in early embryogenesis.

We also observed that the nicked circular biotinylated DNA underwent normal conformational changes into supercoiled DNA. The microinjected heavily biotinylated DNA also underwent efficient replication as indicated by the incorporation of radiolabeled nucleotides (Fig. 5b). We conclude that biotinylated nucleotides can be incorporated efficiently into nascent DNA, are quite stable in early embryos, do not serve as barriers to the synthesis of complementary strands, and do not prevent ligation and torsional stress. The above observations imply that biotinylated nucleotides can be used successfully to study normal replication processes in vivo, both for pulse labeling and for some continuous labeling experiments.

Biotinylation is particularly suitable for studying DNA replication in vivo, because the nascent DNA can be visualized directly both on agarose gels and by electron microscopy. Streptavidin–gold or streptavidin–ferritin conjugates bind efficiently and specifically to the biotinylated portions of the DNA. There is little or no nonspecific binding (Fig. 6). Cross-linking and high backgrounds are eliminated or considerably reduced by the judicious use of biotin-titrated conjugates (Figs. 7 and 8). Our results also show that, although the 20-nm particles gave the highest contrast, smaller particle sizes were more useful for resolving sites of nascent DNA synthesis. Moreover, the large particles impeded spreading of the long molecules of DNA.

The potential resolution of the technique using streptavidin–colloidal gold or streptavidin–ferritin conjugates is such that individual Okazaki fragments (~200 bp) can be identified easily. Two adjacent gold particles with associated streptavidin moieties could theoretically be located at two sites out of phase by 180° and translated along the helical axis by 17.5 Å (Fig. 9). Depending on the size of the gold particle the resolution would be 13, 26, or 50 bp (for 5-, 10-, or 20-nm particles, respectively). In practice, however, technical difficulties involved in spreading dense particle–bound DNA for electron microscopy presumably set a lower limit to the resolution of the method.

The mechanism of chromosomal DNA replication (11–13) in higher eukaryotes has proven elusive (3, 5, 6) during the 35 years since the essential features of duplex DNA replication were first set forth (40). Recently, we have postulated that separation of the strands of the double helix during chromosomal DNA replication in embryos of the frog, Xenopus laevis is temporally and spatially uncoupled from synthesis of the daughter strands (2, 4, 19). The strand separation hypothesis (19) has been proposed also as a possible replication mechanism in human melanoma cells (31a). To distinguish between replication by the strand separation mechanism and replication at processive replication forks we developed the high resolution method presented in this paper to localize sites of nascent DNA synthesis on chromosomal DNA. This method should prove useful also for studying the temporal and spatial relationships of sites of nascent DNA synthesis relative to other cellular processes including, among many others, gene expression and chromatin assembly.

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Figure 8. Streptavidin–colloidal gold conjugate (20-nm size) binding to biotinylated DNA. Streptavidin–colloidal gold was titrated with biotin to optimize binding while minimizing cross-linking and heavy backgrounds. (a) In vitro–biotinylated φX174 Form II DNA. (b) Linearized biotinylated φX174 DNA. (c) In vitro–biotinylated nicked long linear calf thymus DNA labeled with 20-nm colloidal gold. This DNA was relatively sparsely labeled (by nick translation). (d) A short fragment (~1 kb) of heavily biotinylated φX174 DNA reacted with the streptavidin–colloidal gold conjugate. Longer molecules of heavily biotinylated DNA reacted with 20-nm colloidal gold particles do not spread well. (e) DNA as in c, but labeled with streptavidin–ferritin conjugate. Note that binding of ferritin particles does not induce kinks on the spread DNA, unlike the more dense gold particles (b especially).

Figure 9. Computer projection of a representative model of bio-11-dUTP substituted poly(dA)-poly(dU). The figure represents the DNA helix viewed down the helical axis. One turn of the B-DNA duplex with biotin labeled at 5-position on U; the length of the biotin-11 atom linker is ~21–22 Å. Theoretically two gold particles (10–20 nm) with the associated streptavidin moieties could be located at two sites out of phase by 180° and translated along the helical axis by 17.5 Å. Binding of one gold particle (10–20 nm) may protect in principle the whole length of such a segment and stereochemically prevent binding similar particles at many of the available adjacent binding sites. This puts a theoretical constraint on the number of particles that could bind to a certain length of DNA, even though potentially DNA may be biotinylated every eight nucleotides.

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