Expression of β-Nerve Growth Factor and Its Receptor in Rat Seminiferous Epithelium: Specific Function at the Onset of Meiosis

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Abstract. β-Nerve growth factor (NGF) is expressed in spermatogenic cells and has testosterone-downregulated low-affinity receptors on Sertoli cells suggesting a paracrine role in the regulation of spermatogenesis. An analysis of the stage-specific expression of NGF and its low affinity receptor during the cycle of the seminiferous epithelium in the rat revealed NGF mRNA and protein at all stages of the cycle. Tyrosine kinase receptor (trk) mRNA encoding an essential component of the high-affinity NGF receptor was also present at all stages. In contrast, expression of low affinity NGF receptor mRNA was only found in stages VIIα and VIII of the cycle, the sites of onset of meiosis. The low-affinity NGF receptor protein was present in the plasma membrane of the apical Sertoli cell processes as well as in the basal plasma membrane of these cells at stages VIIα to XI. NGF was shown to stimulate in vitro DNA synthesis of seminiferous tubule segments with preleptotene spermatocytes at the onset of meiosis while other segments remained nonresponsive. We conclude that NGF is a meiotic growth factor that acts through Sertoli cells.

β-NERVE growth factor (NGF) is an 118-amino acid protein that is essential for development and maintenance of sensory and sympathetic peripheral neurons (Levi-Montalcini and Angeletti, 1968; Thoenen and Barde, 1980; Levi-Montalcini, 1987). NGF has also been found in the mammalian brain (Korsching et al., 1985; Whittemore et al., 1986; Sheldon and Reichardt, 1986; Goedert et al., 1986) where the factor has been shown to support basal forebrain cholinergic neurons (reviewed in Whittemore and Seiger, 1987; Thoenen et al., 1987; Ebendal, 1989). Recently, NGF has been shown to be a member of a family of structurally related proteins which also includes brain-derived neurotrophic factor (Barde et al., 1982, Leibrock et al., 1989), neurotrophin-3/hippocampus-derived neurotrophic factor (Hohn et al., 1990; Maisonnier et al., 1990; Ernfors et al., 1990; Rosenthal et al., 1990), and neurotrophin-4 (Hallböök et al., 1991).

NGF mediates its effects through interaction with specific receptors (NGF-R) that are present in NGF-responsive neurons of the peripheral and central nervous systems (Banerjee et al., 1973; Herrup and Shooter, 1973; Taniuchi et al., 1986; Richardson et al., 1986; Hefti et al., 1986). The NGF-R can occur in low and high affinity states (Sutter et al., 1979; Landreth and Shooter, 1980; Schechter and Bothwell, 1981), though only the high affinity receptors mediate the biological activity of NGF. Molecular clones have been isolated for the rat, human, and chicken low-affinity NGF receptor (LNGF-R) (Johnson et al., 1986; Radeke et al., 1987; Large et al., 1989). The LNGF-R is a 75-kD glycosylated protein which contains one plasma membrane-spanning domain, a cytoplasmic region, and an extracellular cysteine-rich amino-terminal domain. Chemical cross-linking experiments have demonstrated a NGF-R complex in NGF responsive PC12 cells consisting of the 75-kD low affinity LNGF-R and a second polypeptide of 140-kD (Meakin and Shooter, 1991). The 140-kD component has recently been identified as the trk proto-oncogene, encoding a tyrosine kinase receptor (Kaplan et al., 1991, Klein et al., 1991). Moreover, coexpression of the trk proto-oncogene and the 75-kD low affinity LNGF-R has been shown to result in a high affinity NGF binding postulated to be because of a heterodimer formation between the trk tyrosine kinase receptor and the 75-kD low affinity LNGF-R (Hempstead et al., 1991). The

1. Abbreviations used in this paper: EDS, ethanedimethanesulphonate; LNGF-R, low-affinity NGF receptor; NGF, nerve growth factor; NGF-R, NGF receptor.

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rk proto-oncogene product is a member of a family of at least three structurally related proteins (Martin-Zanca et al., 1989; Klein et al., 1989, 1991; Lamballe et al., 1991). Since brain-derived neurotrophic factor (Rodriguez-Tébar et al., 1990), neurotrophin-3 (Ernfors et al., 1990; Squinto et al., 1991), and neurotrophin-4 (Hallböök et al., 1991) interact with the 75-kD NGF-R with similar binding affinities as NGF, this receptor could be used to mediate the response of all members in the NGF family.

In agreement with a target-derived neurotrophic role of NGF, the levels of NGF mRNA in most peripheral tissues (Heumann et al., 1984; Shelton and Reichardt, 1984) and in the brain (Korsching et al., 1985; Whittome et al., 1986; Shelton and Reichardt, 1986) have been shown to correlate with the degree of innervation by NGF-sensitive fibers. However, in male mouse submandibular gland, snake venom gland (Levi-Montalcini and Angeletti, 1968), guinea pig and rabbit prostate (Harper et al., 1979), and in bull seminal vesicle (Harper et al., 1982), the levels of NGF protein and mRNA do not correlate with the innervation by NGF-sensitive fibers possibly reflecting a non-neurotrophic role of NGF in these tissues. Testis also belongs to such tissues where NGF-like immunoreactivity has been demonstrated in mouse and rat germ cells (Olson et al., 1987; Ayer-Le Lièvre et al., 1988). From these initial data, NGF has been suggested to have a role in sperm maturation and/or motility. In agreement with this possibility, NGF mRNA and protein have been demonstrated in spermatocytes and early spermatids of rat and mouse, and NGF protein has also been detected in the lumen of the epididymis (Ayer-Le Lièvre et al., 1988). Furthermore, the gene for the 75-kD LNGF-R is expressed in Sertoli cells in the testis under negative control of testosterone (Persson et al., 1990) suggesting that NGF produced by spermatogenic cells mediates an interaction between germ and Sertoli cells. Results of in situ hybridizations (Persson et al., 1990) have suggested that the 75-kD LNGF-R is only expressed between stages VI-VIII of the cycle of the seminiferous epithelium (Leblond and Clermont, 1952) providing further evidence for a specific function of NGF and its receptor in spermatogenesis. In this report, we have investigated the functional role of NGF in the testis by accurate localization studies combined with a newly developed in vitro method for studies of stage-specific DNA synthesis during spermatogenesis. Our results show that NGF, presumably by interacting with a stage-specific NGF-R in Sertoli cells, stimulates DNA synthesis at the time of onset of meiosis, suggesting that NGF acts as a meiotic growth factor during spermatogenesis.

Materials and Methods

Seminiferous Tubule Microdissection and Culture

Testes of adult (3-5 mo) Sprague-Dawley rats (ALAB, Sollentuna, Sweden) were decapsulated and seminiferous tubules teased free by fine forceps under transilluminating stereomicroscope in a petri dish containing PBS (Modified Dulbecco's formula, without calcium and magnesium) (Flow Laboratories, Inc., Irvine, Scotland). The stages were recognized according to light absorption criteria (Parvinen and Vainio-Perttula, 1972). For Northern blot analysis, pools of stages I, II-III, IV-V, VI, VIIa-VII, VIIa-VIII, IX-XI, XII, and XIII-XIV, each containing a total of 10 cm of seminiferous tubule segments were collected as described (Parvinen and Ruokonen, 1982). Since LNGF-R is physiologically downregulated by testosterone (Persson et al., 1990), single subcutaneous injection of 75 mg/kg of ethane dimethane sulphonate (EDS) (dissolved in DMSO/water 1:3) was given specifically to destroy the Leydig cells (Jackson et al., 1986). This results in very low levels of intratesticular testosterone for 3-10 d and a marked increase in LNGF receptor and its mRNA (Persson et al., 1990). The same effect can be reached by hypophysectomy. For studies of DNA synthesis in vitro, stages I, V, VIIa, and VIII-IX were prepared from normal adult rat testes (Parvinen et al., 1991). They contain cells at representative phases of mitotic and meiotic DNA synthesis: Stage I, type A-spermatogonia in S phase; stage V, type B-spermatogonia in S phase; VIIa, preleptotene spermatocytes outside S phase; and VIII-IX, preleptotene spermatocytes in S phase (Clermont, 1972). For studies of NGF receptor mRNA expression during in vitro differentiation, quadruplicate pools (2.5 cm each) of stages VIIa and VIIb-VIII were isolated from normal rats for immediate freezing and RNA isolation, or cultured for 48 h at 37 and 34°C in a medium described below before RNA isolation. For immunohistochemical studies, whole testes or ten pools of microdissected 20 x 2 mm seminiferous tubule segments from stages I, II-III, IV-V, VI, VIIa-VII, VIIa-VIII, IX-XI, XII, and XIII-XIV were fixed from normal, hypophysectomized (3 and 10 d), and rats given a single dose of EDS 7 d previously.

RNA Blot-Hybridization Analysis

The frozen seminiferous tubule segments were disrupted in 4 M guanidine isothiocyanate, 0.025 M sodium citrate, pH 7.0, and 0.1 M β-mercaptoethanol followed by centrifugation through a cushion of 5.7 M CsCl in 0.025 M sodium acetate, pH 5.5. After phenol-chloroform extraction, the samples containing 40 μg of RNA were electrophoresed in a 1% agarose gel containing 0.7% formaldehyde. The gel was blotted onto nitrocellulose and hybridized to a 3.2-kb-long insert derived from a rat LNGF-R cDNA clone (Radeke et al., 1987). After washing and exposure to X-ray film, the same filter was boiled for 5 min in 1% glycerol and hybridized to a 456-bp-long polymerase chain reaction fragment from the extracellular region of rat trk. The fragments were labeled with α-32P-dCTP by nick-translation to a specific activity of ~1010 cpm/μg. The filters were washed at high stringency (0.1 x SSC/0.1% NaDodSO4 at 54°C) and exposed to x-ray films at ~80°C with an intensifying screen (DuPont Co., Wilmington, DE).

Immunohistochemical Analyses of NGF

Staged 2-mm tubule segments were microdissected from a normal rat testis. They were fixed in a 4% solution of paraformaldehyde containing 0.2% picric acid for 2 h and cryoprotected with 10% sucrose in PBS. 10 μm cryostat sections were cut with a cryostat (model: HM-500, Micron Instruments, Inc., Los Angeles, CA) and thawed onto gelatin-coated glass slides. The sections were incubated for 12-18 h with a rabbit NGF antibody (K 17, Olson et al., 1987) diluted 1:500 or 1 AM mouse ß-NGF before the immunocytochemical procedure. The sections were incubated with FITC-conjugated goat-anti-rabbit IgG (Boehringer Mannheim, GmbH, Mannheim, Germany) diluted 1:100 for 30 min. The sections were mounted in glycerol/PBS containing 0.1% phenylmedemine and examined with a fluorescent microscope (Nikon FX; Nikon Inc., Melville, NY). The controls included omission of the primary antiserum, replacement of the primary antiserum with normal rabbit serum, and preabsorption of the NGF antiserum with 1 μM mouse β-NGF before the immunocytochemical procedure.

Immunohistochemical Analyses of the 75-kD LNGF-Receptor

Adult control and hypophysectomized (3 and 10 d) male rats were perfused transcardially with the same fixative as above. The testes were then excised and immersion fixed for 2 h. Staged 2 mm tubule segments from EDS-treated rats (7 d after a single dose) were treated as described above. The sections were incubated overnight with mouse monoclonal rat 75-kD LNGF-R antibody (192-IgG, Taníuchi and Johnson, 1985) diluted 1:500 or 1:1,000 in PBS containing 1% BSA and 0.3% Triton X-100. Subsequently, the sections were incubated with FITC-conjugated goat-anti-rabbit IgG (Boehringer Mannheim, GmbH, Mannheim, Germany) diluted 1:100 for 30 min. The sections were mounted in glycerol/PBS containing 0.1% phenylmedemine and examined with a fluorescent microscope (Nikon FX; Nikon Inc., Melville, NY). The controls included omission of the primary antiserum, replacement of the primary antiserum with normal mouse serum (1:500). Diaminobenzidine was used as a chromogen.
removed and immersion fixed for 6–12 h. The tissues were cryoprotected with 50% sucrose in PBS for several days and frozen in liquid nitrogen. Thick sections (~500 µm) were cut with a razor blade and allowed to thaw in PBS. The sections were then incubated with mouse mAb to the 75-kD LNGF-R diluted 1:500 for 72 h. After several washes, the tissues were incubated in biotinylated goat-anti-mouse IgG and ABC-complex for 2 h each. Diaminobenzidine was used as chromogen to reveal ABC-complexes. Subsequently, the tissues were postfixed with 2.5% glutaraldehyde, 1% osmium tetroxide, and 1% uranyl acetate for 30 min each. The tissues were dehydrated and flat embedded in Epon. Sections exhibiting immunolabeling were processed for EM and the ultrastructural examination was undertaken with a microscope (model 1200EX; JEOL USA, Inc., Peabody, MA) without counterstaining.

Analysis of DNA-Synthesis
The staged 2-mm tubular segments from normal rats were individually transferred in 10 µl of PBS into 96-well tissue culture plates (Falcon 3072 Microtest III; Becton Dickinson Microbiology Sys., Cockeysville, MD) and incubated for 24, 48, or 72 h in 100 µl alpha modification of MEM (α-MEM) with Earl's salts (12:312:54; Flow Laboratories, Inc.) supplemented with L-glutamine (2 mM), penicillin (100 U/ml), streptomycin (100 µg/ml), and α-thioglycerol (7.5 × 10⁻² M) (Sigma Chemical Co., St. Louis, MO). During the last 4 h, 20 kBq of methyl-³H]thymidine (TRA.120, 185 GBq/mmol, Amersham International, Amersham, UK) was added.

Effects of β-NGF on [³H]Thymidine Incorporation
After microdissection of 2-mm segments of stages I, V, VII, and VIII–IX, they were transferred to tissue culture wells as described above and incubated for 24, 48, or 72 h in the presence of 0 (control), 40, 100, and 250 ng/ml of β-NGF purified from the male mouse submandibular gland (Ebendaal et al., 1984). Cultures were maintained in α-MEM supplemented as described above at a temperature of 37 or 34°C (physiological) in a humidified atmosphere of 5% CO₂ in air. Three separate experiments were performed, each with six replicate samples.

NGF Preparation and Purification
β-NGF was purified from male mouse (NMRI, body weight >30 g) submandibular glands as previously described (Ebendaal et al., 1984). Elution from the second column of carboxymethyl-Sepharose was by a linear gradient of sodium chloride yielding a peak of highly purified β-NGF (Ebendaal et al., 1984; Olson et al., 1991). The concentration of NGF was determined from specific absorbance at 280 nm (1.6 at 1 mg/ml in a 1-cm cuvette). Identity and purification was further verified by electrophoresis in sodium dodecylsulphate on a gradient polyacrylamide gel, and by immunoblotting of similar gels (Olson et al., 1991).

Bioassay of NGF Activity
The biological activity of NGF was measured in a nerve fiber outgrowth assay with explanted sympathetic ganglia from the 9-d-old chick embryo. The dissected ganglia were explanted into a collagen matrix (Ebendaal et al., 1984). To determine activity, the purified mouse NGF was added in a series of twofold dilutions with final concentrations ranging from 0 to 20 ng/ml. The cultures were incubated at 37°C using 5% CO₂ in air. Three separate experiments were performed, each with six replicates samples.

Cell Harvesting and Radioactivity Measurement
After [³H]thymidine labeling, the tissue cultures were harvested by a Titerhek Cell Harvester 550 (Flow Laboratories, Inc.) for 1 min/row using distilled water and Titerhek glass-fiber filter paper (Skatron, Inc., Lier, Norway). The filter disks were transferred into minivials by a punching apparatus (Skatron, Inc.) for liquid scintillation counting by a spectrometer utilizing Ready Safe (Model LS500CE, Beckman Instruments, Inc., Fullerton, CA) scintillation cocktail.
H., and J.-P. Merlio, unpublished results). In control animals not treated with EDS, the 75-kD LNGF-R mRNA expression at stages VII_a–VIII was much lower, but a clear difference to stages VII_b could be demonstrated (Fig. 4). When seminiferous tubular segments from stages VII_a and VII_b–VIII were cultured in chemically defined serum-free medium for 48 h, low levels of 75-kD LNGF-R mRNA expression appeared in tubular segments which were at stages VII_a at the onset of culture suggesting that a differentiation occurred in vitro. This differentiation only occurred at the physiological temperature of 34°C; no 75-kD LNGF-R mRNA signal was detected when the seminiferous tubular segments were cultured at 37°C (Fig. 4).

In immunohistochemical analysis of staged seminiferous tubule segments from EDS-treated rats using an mAb specific for the 75-kD LNGF-R, some of the tubules at stages VII_a showed a reaction in the apical parts of the Sertoli cells. In contrast, all the tubules at stages VII_b and VIII showed 75-kD LNGF-R immunoreactivity mainly at the apical portions of the Sertoli cells (Fig. 5). Some reaction was also

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**Figure 1.** (A) Fluorescent immunocytochemical analysis of the expression of NGF protein during spermatogenesis (stages of the cycle are indicated at lower left corner of each panel). The spermatogenic cells in all stages of the cycle showed immunoreactivity. In stages I–V, mainly round spermatids were labeled; in stages VI–VIII both round spermatids and pachytene spermatocytes; and in stages IX–XIV mainly pachytene, diakinetic, and dividing spermatocytes were labeled. During stages IX–XIV, a declining immunoreactivity was seen in acrosome phase spermatids (steps 9–14). Spermatogonia and early spermatocytes for up to mid-pachytene as well as maturation phase spermatids (steps 15–19) were negative. (B) A stage IV–V control tubule incubated with an antiserum preabsorbed with NGF. No specific NGF immunostaining is seen in the layers of spermatogonia (G), spermatocytes (C), round spermatids (R), or elongated spermatids (E). The finding was similar in all stages of the cycle. Bars, 25 μm.

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**Figure 2.** Northern blot analysis of a 75-kD LNGF-R mRNA in seminiferous tubule segments microdissected from defined stages of the epithelial cycle 4 d after a single dose of EDS. Total cellular RNA (40 μg/slot) was electrophoresed in a formaldehyde containing agarose gel, blotted onto a nitrocellulose filter, and hybridized to a cDNA probe for the 75-kD low-affinity NGF-R. The filter was washed at high stringency and exposed to x-ray film.

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**Figure 3.** Northern blot analysis of rat *trk* mRNA in the same filter as in Fig. 2.
progressed at a slightly accelerated rate as compared with the physiological temperature of the testis. At 37°C, spermatogenesis that observed in vivo (Clermont and Harvey, 1965). After 24 h at 37°C, the highest basal DNA synthesis was observed in seminiferous tubules from stages VIII-IX of the cycle (Fig. 9 A). A significant dose-dependent stimulation of premeiotic DNA synthesis was found after a 48-h culture. All concentrations of NGF had a stimulatory effect on what was originally stage VII, tubular segments at their onset of meiotic DNA synthesis (Fig. 9 B). The autoradiographic analysis revealed that the cells incorporating radioactivity in these conditions were preleptotene spermatocytes at the onset of meiosis (Fig. 10). The other stages showed no response. At 72 h, the meiotic DNA synthesis at stage VII was fully activated, and the stimulatory or maintaining effects of NGF were no longer seen (Fig. 9 C).

Discussion

Fig. 11 summarizes the findings of the present work and relates them to the previously described stage-specific DNA synthesis during spermatogenesis (Parvinen et al., 1991). LNGF protein is expressed in spermatogenic cells from late meiosis to the mid-spermiogenesis, but its receptor is only expressed in Sertoli cells at stages VIIa and VIII where preleptotene spermatocytes activate in the meiotic DNA synthesis. NGF appears to be the first growth factor described that specifically affects meiosis by mediating an interaction between germ cells and Sertoli cells. Although NGF is known to have important functions in the development and maintenance of sensory and sympathetic neurons in the mammalian nervous system (Levi-Montalcini, 1987), its function in the testis has remained obscure. However, the strictly limited expression of 75-kD LNGF receptor mRNA to Sertoli cells at stages VIIa and VIII of the seminiferous epithelial cycle suggests a specific paracrine function. In contrast, NGF mRNA and protein did not show such a stage-specific localization and were found in primary spermatocytes and in spermatids at all stages. The expression of trk mRNA did not show a great variation during the cycle suggesting that the trk proto-oncogene protein is present in the testis although its expression is not regulated by the same strictly stage-specific manner as that of the 75-kD LNGF-R. Provided that the 75-kD LNGF-R can mediate a response of NGF in the testis, perhaps as a heterodimer between the 75-kD LNGF-R and trk (Hempstead et al., 1991), events at stages VIIa and VIII of the cycle may be targets for NGF-mediated actions.

Stages VII and VIII of the cycle of the seminiferous epithelium are preferentially regulated by androgens (Parvinen, 1982), and synthesis of specific proteins in these stages is influenced by testosterone (Sharpe et al., 1992) and by germ cells (McKinnell and Sharpe, 1992). Surprisingly, the ex-
Figure 5. Immunohistochemical analysis of 75-kD LNGF-R protein during the cycle of rat seminiferous epithelium in a rat treated with EDS 7 d previously (stages are indicated at lower left corner of each panel). Some of the tubules at stage VIIb showed a reaction in the apical parts of the Sertoli cells, whereas all the tubules at stages VIIc and VIII showed immunoreactivity mainly at the apical portions of the Sertoli cells. An increasing reactivity was found at the basal part of Sertoli cells during stages VIIc-XI (arrows). In stage XII, the reactivity was disappearing. Stages XIII-VI were negative. Bar, 50 μm.
pression of 75-kD LNGF-R mRNA and protein is regulated in a negative fashion by testosterone at these stages. This further supports the central role of androgens in the regulation of this part of the seminiferous epithelium. In terms of DNA synthesis, the same stages belong to the most active ones during the cycle of the seminiferous epithelium. The A₁ spermatogonia located most basally in the seminiferous epithelium enter S phase during stages VIIa-VIII and undergo the first mitotic division that starts the entire spermatogenesis (Clermont, 1972). However, the A₁-spermatogonia are too few to significantly contribute to the total DNA synthesis of stages VIIa-VIII. The preleptotene spermatocytes that enter DNA synthesis to start meiosis at the same stage of the cycle are far more abundant in number (Wing and Christensen, 1982). Without doubt they contribute the bulk of [³H]thymidine incorporation considered as meiotic DNA synthesis, also revealed by autoradiography (Monesi, 1962; Parvinen et al., 1991). Spermatocytes at the onset of meiosis are displaced from the basal lamina by Sertoli cell projections (Russell, 1977) in a process suggested to be regulated by plasminogen activator (Lacroix et al., 1981). In fact, secretion of plasminogen activator is maximal at the border between stages VII and VIII of the cycle when analyzed in sequentially cut 1 mm seminiferous tubule segments (Vihko et al., 1984). This distribution resembles that of the 75-kD LNGF-R mRNA. At stages VII and VIII of the cycle, the pachytene stage primary spermatocytes are maximally activated in their meiotic RNA synthesis but step eight spermatids cease their transcriptional activity (Monesi, 1965; Söderström and Parvinen, 1976). Finally, the most mature step 19 spermatids become released from the seminiferous epithelium at stage VIII of the epithelial cycle (Leblond and Clermont, 1952). The present data do not allow us to evaluate the role of NGF in the regulation of all of these processes. However, a hypothesis to explain the existence of the constant cell associations in the seminiferous epithelium is that different generations of spermatogenic cells require similar metabolic activation of the Sertoli cells. This may be the case for DNA synthesis in A₁-spermatogonia and preleptotene spermatocytes at the same stage of the seminiferous epithelial cycle.

The new method for quantitative analysis of DNA synthesis during rat spermatogenesis (Parvinen et al., 1991) is sometimes influenced by unexpected modulations of the seminiferous epithelial wave (Perey et al., 1961), particularly in rats that were more than 5-mo-old. Therefore, only rats between 60 and 90 d of age gave results with a variability small enough to reveal the subtle effects of NGF. In all positive experimental conditions (four out of six), the highest concentration of NGF significantly stimulated the premei-
Figure 7. Electron microscopic immunohistochemical analysis of 75-kD LNGF-R protein 10 d after hypophysectomy. (A) The main reactivity was found in the apical Sertoli cell processes approximately at stages VII<sub>a</sub>-VIII of the cycle (arrow); the adjacent germ cells were negative (A). (B) The immunoreactivity was associated with plasma membranes and it was particularly strong in the narrow and long extensions of the Sertoli cells (arrows). (C) 75-kD LNGF-R immunoreactivity was also seen in the basal plasma membrane of the Sertoli cell (arrows) around the typical indented nucleus (n). (D) Some Sertoli cells showed immunoreactivities only at limited areas of the plasma membrane (arrows); m, mitochondrion typical for Sertoli cell. Bars, 500 nm.
Figure 8. In vitro DNA synthesis (cpm incorporated from $[^3H]$thymidine during the last 4 h of culture) of staged 2-mm seminiferous tubule segments after 24 (A), 48 (B), and 72 (C) h at 37°C (means ± SEM, n = 18). The approximate progression of stages V, VIIa, VIII–IX, and I of the seminiferous epithelial cycle is indicated by arrows at the x-axis of each panel. After 24 h, the preleptotene–leptotene spermatocytes at stage X are still active in DNA synthesis, and NGF has a dose-dependent maintaining effect on this. After 48 h, the resting preleptotene spermatocytes at original stage VIIa have spontaneously activated in meiotic DNA synthesis (B), but NGF had no effect. However, after 72 h, when stage VIIa has differentiated to IX, preleptotene spermatocytes are activated in meiotic DNA synthesis and NGF has a dose-dependent effect (C). Premiotic DNA synthesis was not stimulated or maintained by NGF in any conditions examined. *, $P < 0.05; **$, $P < 0.01$ compared with controls.

Figure 9. In vitro DNA synthesis (cpm incorporated from $[^3H]$thymidine during the last 4 h of culture) of staged 2-mm seminiferous tubule segments after 24 (A), 48 (B) and 72 (C) h at physiological temperature 34°C (means ± SEM, n = 18 except stage VIIa after 48 h, where n = 14). The approximate progression (according to Clermont and Harvey, 1965) of each stage is indicated by arrows at x-axes. After 24 h, the preleptotene–leptotene spermatocytes of stage IX are still active in DNA synthesis, and NGF has a slight dose-dependent stimulating effect (A). After 48 h, most of the preleptotene spermatocytes of stage VIIa tubule segments have not yet spontaneously activated in meiotic DNA synthesis, but NGF significantly enhances this activation (B). After 72 h, preleptotene spermatocytes of original stage VIIa had spontaneously activated in DNA synthesis (C), but NGF had no effect. Premiotic DNA synthesis was not stimulated or maintained by NGF in any conditions examined. *, $P < 0.05; **$, $P < 0.01$ compared with controls.

The absence of NGF effect in two conditions (48 h/37°C and 72 h/34°C) is puzzling and may be because of an interaction of growth promoters and thus far poorly investigated inhibitors of DNA synthesis. The physiological temperature of the testis (34°C) seems to be important and NGF clearly had a promoting effect on the premeiotic DNA synthesis. It is possible that in adverse conditions such as cryptorchidism, NGF is particularly important in maintaining and supporting the testicular functions. In agreement with this, a stabilizing effect of NGF has been demonstrated...
in cultures of human seminiferous tubules (Seidl and Holstein, 1990a,b).

The absence of any effect on DNA synthesis at stages I and V containing spermatogonia in S phase, but a dose-dependent stimulation at stages VII and VIII-IX shows that NGF in vitro stimulates DNA synthesis at the onset of meiosis but not during mitotic proliferation of spermatogonia. Our data suggest that NGF synthesized in male germ cells stimulates meiosis through interaction with testosterone regulated stage-specific receptor on Sertoli cells. Other

Figure 11. Summary of the findings of the present study superimposed on the map of spermatogenesis (modified from Perey et al., 1961). The stage-specific DNA-synthesis ( cpn x 10^-3/mm, Parvinen et al., 1991) has two major peaks, at stages V (premitotic) and VIII (premeiotic). LNGF protein is expressed in spermatogenic cells from stage VI pachytene spermatocytes to step 14 spermatids ( ), whereas pachytene spermatocytes ( ), round spermatids ( ) and elongated spermatids ( ) remained unlabelled. Bar, 25 μm.

Figure 10. [3H]thymidine (5 kBq/well) autoradiography of a squash preparation of stage VII, tubule segment cultured for 48 h at 34°C in the presence of 250 ng/ml NGF. The preleptotene spermatocytes with small dark nuclei are labeled (arrow), whereas pachytene spermatocytes (P), round spermatids (R) and elongated spermatids (E) remained unlabelled. Bar, 25 μm.
growth factors may have functions in regulating S phases of type A2-4, intermediate and type B-spermatogonia. Interleukin 1α is an example of a testicular growth factor that has effects on both spermatogonial (mitotic) and meiotic DNA synthesis (Parvinen et al., 1991), and there is evidence that IGF-1 is not mitotic but only a spermatogonial growth factor (Söder et al. 1991). It appears that a very complex interaction of several growth factors together with follicle stimulating hormone and testosterone ultimately leads to a proper regulation of spermatogenesis.

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