STATHMIN EXPRESSION MODULATES MIGRATORY PROPERTIES OF GN-11 NEURONS IN VITRO.


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Abstract

Expression of stathmin, a microtubule-associated cytoplasmic protein, prominently localized in neuroproliferative zones and neuronal migration pathways in brain, was investigated in the GnRH neuroendocrine system in vivo and the function was analysed using an in vitro approach.

Here we present novel data demonstrating that GnRH migrating neurons in nasal regions and basal forebrain areas of mouse embryos express stathmin protein. In addition, this expression pattern is dependent on location, as GnRH neurons reaching the hypothalamus are stathmin-negative. Immortalized GN-11 cells, which retain many characteristics of migrating GnRH neurons, strongly express stathmin mRNA and protein. The role of stathmin in GnRH migratory properties was evaluated using GN-11 cell line. We up- (STMN+) and down-regulated (STMN-), the expression of stathmin in GN-11 cells and we investigated changes in cell morphology and motility in vitro. Cells over-expressing stathmin assume a spindle-shaped morphology and their proliferation, as well as their motility, is higher with respect to parental cells. Furthermore, they do not aggregate and express low levels of cadherins as compared to control cells. STMN- GN-11 cells, are endowed with multipolar processes, they show a decreased motility and express high levels of cadherin protein. Our findings suggest that stathmin plays a permissive role in GnRH cell motility, possibly via modulation of cadherins expression.

Abbreviations: STMN: stathmin transfected clones
               CNS: central nervous system
               DMEM: Dulbecco's Modified Eagle Medium
               FBS: foetal bovine serum
               GAPDH: enzyme glyceraldehyde-3-phosphate dehydrogenase
GnRH: Gonadotropin-releasing hormone
HGF/SF: Hepatocyte Growth Factor / Scatter Factor
MT: microtubule
PAF: paraformaldehyde
PBS: phosphate-buffered saline
SFM: Serum Free Medium
Introduction

Gonadotropin-releasing hormone neurons (GnRH, also known as luteinizing hormone-releasing hormone, LHRH) control the release of anterior pituitary hormones, LH and FSH (follicle stimulating hormone), modulating the reproductive functions (1). In adult vertebrates these neuroendocrine cells are scattered throughout the ventral forebrain and reside predominantly in the septal-preoptic and anterior hypothalamic areas (2). During embryonic development, GnRH cells originate in the nasal placode and migrate along olfactory/vomeronasal pathway to their final locations into the brain (3). Although the migration of these cells, from nose to brain, has been well documented in a variety of species (4, 5, 6, 7, 8, 9, 10), many questions remain concerning the molecules and cues directing GnRH cell migration.

Cell migration is a complex cellular behaviour that results from the coordinated changes in the cytoskeleton and the controlled formation and dispersal of adhesion sites. Whereas the actin cytoskeleton provides the driving force at the cell front, the microtubule (MT) network assumes a regulatory function in coordinating rear retraction (11). Cytoskeletal rearrangements are crucial for all these events. For example, leading edge extension depends on polymerization of actin microfilaments, and nuclear translocation and retraction of the trailing edge involve MT assembly and disassembly, respectively (12). Stathmin (13) is a member of a class of MT destabilizing proteins that regulate the dynamics of MT polymerization/depolymerization and plays a critical role in the regulation of the dynamic equilibrium of MTs during different phases of the cell cycle (14; 15; 16; 17; 18; 19) in a phosphorylation-dependent manner (14; 16; 20). The MT destabilizing activity of stathmin is turned off by cell surface receptor kinase cascades and cycle-dependent kinases (21; 22; 23; 24; 25).
In the adult nervous system of mammals, relevant neurogenesis occurs in restricted areas, such as the olfactory epithelium (OE), the subependymal layer and the dentate gyrus of the hippocampus (26; 27). Previous studies have demonstrated a high level of stathmin expression in these regions, supporting the idea that this protein is involved in processes of migration and differentiation (28). Interestingly, stathmin expression has been documented in the developing main olfactory epithelium and vomeronasal organ during mouse embryonic development as well as in populations of migrating cells crossing the nasal mesenchyme at early stages of development and resembling, for morphology and location, GnRH neurons (29). Here we show that stathmin is expressed in migrating GnRH neurons in vivo and in immortalized highly migratory GnRH neurons (GN-11 cells). Furthermore, using stable transfected GN-11 cells in which stathmin was up- or down-regulated we show that stathmin specifically modulates motility properties of GN-11 cells.
Materials and methods

All reagents were purchased from SIGMA-ALDRICH, St. Louis, Missouri, USA unless otherwise specified.

Animals and tissue

The experiments were carried out on CD-1 mice (Charles River, Lecco, Italy). All animal protocols were approved by the Animal Care and Use Committee of the University of Turin.

Embryos

Timed pregnant mice (plug day, E0) were deeply anaesthetized and sacrificed at specified embryonic days. Embryos were harvested and washed in ice-cold Dulbecco's phosphate buffer (PBS). The heads were fixed in fresh 0.1 M pH 7.4 PBS, 4% paraformaldehyde (PAF) overnight at 4°C and cryoprotected in sucrose solution the next day. Tissues were then embedded in Killik frozen section medium (Bio Optic, Milan, Italy) and frozen in liquid nitrogen-cooled isopentane. Sagittal sections were cut, mounted on 3-aminopropyl-trietoxysilane-treated slides (Fluka, Milan, Italy) and stored at –20°C until processing for immunohistochemistry.

Adult mice

Adult CD-1 mice postnatal day 30 (P30) were anaesthetized with an intraperitoneal injection of Ketamine (200 mg/kg), and perfused with 4% PAF. The brains were dissected and postfixed in the same fixative overnight at 4°C, cryoprotected in sucrose solutions and sectioned on a cryostat. Sagittal sections were cut, mounted on 3-aminopropyl-trietoxysilane-treated slides (Fluka, Milan, Italy) and stored at –20°C until processing for immunohistochemistry.
8 μm thick sections were used for consecutive mirror reactions, 14 μm thick sections for simultaneous double-labelling reactions.

**Cell cultures**

GN-11 cells (30) were grown in monolayer at 37 °C in a humidified atmosphere of 5% CO₂/air, in Dulbecco's modified Eagle's medium (DMEM 4500 mg glucose) containing 1 mM sodium pyruvate, 2 mM glutamine, 100 μg/ml streptomycin, 100 U/ml penicillin and supplemented with 10% Foetal Bovine Serum (FBS, Invitrogen, Inc., Grand Island, NY). The medium was replaced at 2-day intervals. Subconfluent cells were routinely harvested by trypsinization and seeded in 57 cm² dishes (1 x 10⁶ cells).

**Plasmids construction**

DNA manipulations were carried out using standard techniques (31). cDNA fragments containing the entire coding region of rat stathmin were amplified by RT-PCR. The primers used (produced by Sigma-Genosys) were designed according to the rat GenBank® stathmin sequence (accession number J04979): 5’-TGTCTTCTGTCCAACATGGC-3’ and 5’-AAAACATCTCACGGTCTGGA-3’ corresponding to nucleotides 154-766. The obtained fragments were purified from agarose gel using GenElute™ Gel Purification Kit and cloned into the pGEM®-T Vector System (Promega Corporation, Madison, WI, USA) following the manufacturer's instructions; the target DNA amplification fragment was then sequenced. The DNA of single colonies was obtained using GenElute™ Plasmid Miniprep Kit and subcloned into the EcoR V gap of pIRESpuro2 Vector (CLONTECH Laboratories, Palo Alto, CA). Stathmin antisense and stathmin sense expression vectors were identified by Xbal restriction map and denoted respectively pSTATHMIN- and pSTATHMIN+. 
Plasmid transfection and generation of stable cell lines

pSTATHMIN-, pSTATHMIN+ and pIRESpuro2 empty vector, as a control, were transfected in GN-11 cells. Transfections were performed using Lipofectamine™ Reagent transfection reagent (Invitrogen™ Life Technologies, Inc., Grand Island, NY) and Optimem (Invitrogen™ Life Technologies, Inc., Grand Island, NY) according to the manufacturer's instruction. Briefly, cells were seeded in 100-mm tissue culture plates one day prior to transfection. The cell cultures, 60-80% confluent, were transfected with 10 µg plasmid DNA per plate for 4 hours. After 48 hours incubation in the appropriate growth medium, the selection of clones was carried out for 2 weeks in 10 µg/ml puromycin. About thirty different clones were collected and assayed for stathmin expression; clones expressing high or low stathmin levels were denoted respectively STMN+ or STMN- cells.

Immunohistochemistry

Consecutive E12 and P30 mouse sagittal sections (8 µm) were stained for GnRH and stathmin immunoreactivity. GnRH neurons were labelled using a rabbit polyclonal anti-GnRH antibody (LR5; a generous gift from Dr. Benoit, Montreal, Canada) at a 1:10000 dilution, whereas stathmin immunoreactivity was detected using a rabbit polyclonal stathmin antiserum (a generous gift from Dr. Sobel, Paris, France) at a 1:1000 dilution. Sections were incubated overnight at 4°C with the mentioned antibodies diluted in 1% normal goat serum and 0.3% triton X-100 in PBS. Sections were then washed in PBS and incubated with anti-rabbit biotinylated secondary antiserum and the Vector kit (ABC kit, Vector Laboratories, Burlingame, CA, USA) and processed for avidin-biotin-horseradish peroxidase/3'-3'-diaminobenzidine (DAB) histochemistry. Sections were then washed in PBS, mounted and cover slipped. Negative controls were run by omitting the primary antibody.
E12, E18 and P30 mouse sagittal sections (14 µm) were double-labelled for GnRH and stathmin immunoreactivity. Sections were incubated overnight at 4°C with the mentioned antibodies diluted in 1% normal goat serum and 0.3% triton X-100 in PBS. The following day, sections were washed in PBS and incubated for 1h with anti-Rb-Cy2-conjugated antibody (1:500, Jackson Immunoresearch, West Grove, PA), washed in PBS, mounted and cover slipped. For double-labelling reactions anti-GnRH/anti-stathmin, the AffiniPure Fab Fragment Goat Anti-Rabbit IgG (H+L) (FabBlocker, Jackson Immunoresearch, West Grove, PA) was used according to the manufacture protocol.

Labelled sections were mounted, air-dried, and cover slipped in polyvinyl alcohol with the anti-fading mounting medium diazabicyclo-octane (DABCO). Fluorescent signals were detected using a FV 200 Olympus Fluoview confocal laser scanning microscope. Only adjustment to brightness and contrast were used in the preparation of the figures.

For immunocytochemical analysis of STMN mutants, cells were grown on glass dishes in 10% FBS DMEM. After 24 hours cells were washed twice with PBS, and fixed for 6 minutes in 100% methanol at -20°C. Microtubule networks were visualized using the α-tubulin antibody (mouse monoclonal 1:1000, SIGMA-ALDRICH, St. Louis, Missouri, USA) and the anti-Ms-Cy3-conjugated antibody (1:500, Jackson Immunoresearch, West Grove, PA). Labelled cells were observed on an IX50 inverted microscope (Olimpus Corp., Hamburg, Germany) equipped with a CCD camera CoolSNAP-Pro (Media Cybernetics, Huston, TX) and images edited with Image Pro-Plus software (Media Cybernetics, Huston, TX).

**RT-PCR analysis**

Total RNA was isolated by extraction with TRIzol (Invitrogen™ Life technologies, Inc., Grand Island, NY). Single strand cDNA was synthesized by M-MLV reverse transcriptase from 1µg of total RNA primed with 50 pmol of random hexamers (Amersham
Biosciences, Piscataway, NJ) in a 20 µl reaction. Each reaction consisted of Gibco’s first strand cDNA synthesis buffer (50 mM Tris-HCl pH 8.3, 75 mM KCl, 3 mM MgCl₂), 4 mM dNTPs (Amersham Biosciences, Piscataway, NJ), 1.8 U/µl RNAsin (Amersham Biosciences, Piscataway, NJ) and 10 U/µl M-MLV reverse transcriptase (Invitrogen™ Life technologies, Inc., Grand Island, NY). Samples were incubated at 37°C for 1h. Negative controls: (RT-) were those in which reverse transcriptase was not added, (H₂O) those in which sterile water instead of cDNA was added.

PCR was carried out using 5 µl of cDNA and the appropriate oligonucleotides (0.6-6 µM) in a 30 µl PCR reaction using standard reaction buffer (10 mM Tris-HCl pH 8.3, 50 mM KCl, 1.5 mM MgCl₂, 0.001% gelatin), 0.8 mM dNTPs (Amersham Biosciences, Piscataway, NJ) and 0.025 U/µl of REDTaq DNA polymerase.

The following primers were used: 5’-TGGCATTGTGGAAGGGCTCATGAC-3’ and 5’-ATGCCAGTGAGCTTCCCGTTCAGC-3’ for GAPDH amplification corresponding to nucleotides 544-732 (accession number M32599); 5’-TGTCTTCTGTCCAACATGGC-3’ and 5’-AAAACATCTCACGGTCTGGA-3’ for stathmin amplification corresponding to nucleotides 154-766 (accession number J04979); 5’-GGGACTGCAGCAGCAAAGC-3’ and 5’-GTCTGAGCATCTAGAGTTTCC-3’ for c-Met amplification corresponding to nucleotides 296-815 (32). The amplification of GAPDH served as a control with respect to the quality and quantity of RNA that had been retro-transcribed into cDNA. The number of cycles and the annealing temperature used for each primer pair were: 30 cycles at 57°C for stathmin, 25 cycles at 62°C for c-Met and 25 cycles at 60°C for GAPDH. Amplification products were separated by 1.5% agarose gel electrophoresis and DNA bands visualized by ethidium bromide staining.

*Western blot analysis*
For western-blotting analysis, GN-11 cells were solubilized in lysis buffer (Tris-HCl pH 7.4, 100 mM NaCl, 5 mM MgCl₂, 0.5% NaDOC, 1% Nonidet P40, 1 mM PMSF, 2 mM orthovanadate) on ice. Lysates were clarified by centrifugation at 14000 g for 15 min and protein content determined using a bicinchoninic acid kit for protein determination. Equal amounts of proteins were boiled in Laemmli buffer (2% SDS, 50mM Tris-HCl pH 7.4, 20% β-mercaptoethanol, 20% glycerol) and subjected to 8% (cadherins and c-Met) and 15% (stathmin) SDS polyacrylamide gel electrophoresis (PAGE). Proteins were blotted onto Hybond-C Extra membrane (Amersham Biosciences, Piscataway, NJ).

After blocking with 5% dry milk in TBST buffer (20 mM Tris, 150 mM NaCl, 0.1% Tween 20, pH 7.4) filters were probed with 1:500 monoclonal ANTI-PAN cadherin antiserum, or 1:500 polyclonal c-Met antiserum (Santa Cruz Biotechnology, Inc.), or 1:10000 polyclonal anti-stathmin antibody, or 1:1000 monoclonal β-actin antiserum (SIGMA-ALDRICH, St. Louis, Missouri, USA) and visualized with the appropriate peroxidase-coupled secondary antibodies using ECL detection system (Amersham Biosciences, Piscataway, NJ).

**Proliferation assay**

Cellular proliferation was quantified with the *96-well plates technique* as previously described (33; 34). Briefly, cells were plated in 200 µl serum free medium (SFM) at a density of 1000 cells/well in 96-well plates. The following day (t=0) the media was replaced by 100 µl/well SFM with or without 50 ng/ml of human recombinant HGF/SF; for each treatment, 8 wells were used. At t=0 a 96-well plate was fixed and used as a cell growth start point. Cell growth was subsequently calculated at 1 (t=24h), 2 (t=48h) and 3 (t=96h) days after the initiation of treatments. The medium was removed and cells were fixed by addition of 100 µl 2% glutaraldehyde in PBS. After being shaken (200 cycles/min) for 20 min at room temperature, plates were washed five times by submersion in deionised water.
and air dried at least 24 h. Plates were then stained by addition of 100 µl/well of a solution containing 0.1% crystal violet dissolved in freshly prepared 200 mM boric acid at pH 9.0. After being shaken (200 cycles/min) for 20 min at room temperature, plates were washed five times by submersion in deionised water and air dried at least 24 h. Bound dye was solubilized by addition of 100µl/well of 10% acetic acid and 5 min shaking at room temperature. The optical density of dye extracts was measured directly in plates using a Microplate Reader (BIORAD), at the wavelength of 590 nm.

**Morphological analysis of stable transfected clones**

In order to classify and analyse different morphologies displayed by STMN+, pIRES and STMN- cells, Image Pro-Plus (Media Cybernetics, Silver Spring) and Photoshop 7.0 softwares were used. Random cells were photographed using a 1X50 inverted microscope (Olimpus Corp., Hamburg, Germany) equipped with a CCD camera CoolSNAP-Pro (Media Cybernetics, Houston, TX). For morphometric analysis cell body boundaries were manually traced with processes extending from cell bodies being not taken into account for measurements. Area, perimeter, maximum and minimum diameter of each cell were calculated using the software Image Pro-Plus (Media Cybernetics, Silver Spring). Moreover, for each cell the ratio between maximum and minimum diameter of the cell body was evaluated. The measures were obtained considering only interphasic cells. All data were collected and statistically analysed.

**Motility assay**

To measure two-dimensional movement, cells were plated at low density in culture dishes. The day after plating, cells (in DMEM-10%FBS) were photographed every 30 minutes for 3 hours and motility was analysed as nuclear movement measurements.
Photographs were aligned using Reconstruct Software and nuclear coordinates were measured using ImageJ Software. All the data were collected and statistically analysed.

**Aggregation assay**

Cells aggregates were prepared by the *hanging drop technique* (35). Subconfluent cells were collected by trypsinization, resuspended in complete culture medium and $20^5$ cells were seeded in 20 µl drops on the lid of a culture dish; the lid was then placed on a 35 mm dish filled with 2 ml of culture medium and incubated at 37 °C for 48 h.

Collagen gel was obtained as previously described (36) and 200 µl were pipetted onto the bottom of a well of a 24-well culture dish, and left to set at room temperature. Cells aggregates were transferred over the cushion and then overlaid with additional 200 µl of collagen and covered with culture medium; aggregates were then photographed after 1 hour.

**Migration assay**

The Transwell migration assay was used to measure three-dimensional movement. $10^5$ cells were seeded on the upper side of a FALCON™ cell culture insert (Becton Dickinson Labware, Franklin Lakes,NJ) on a porous polycarbonate membrane (8 µm pore size, 1x 105 pores/cm²) (37); the lower chamber of the cell plate was filled with serum-free DMEM or with serum-free DMEM containing 50 ng/ml of human recombinant HGF/SF. After 6 hours of incubation, cells attached to the upper side of the filter were mechanically removed, whereas cells that migrated to the lower side of the filter were fixed in PBS 1% glutaraldehyde, stained with toluidine blue, photographed (4mm²/ sample) and counted using Image Pro-Plus software (Media Cybernetics, Silver Spring).
Statistical analysis

The experiments were performed in triplicates, and all counts obtained from assays were analysed, averaged and expressed as mean ± standard error. SPSS 12.0 software (SPSS, Inc., Chicago, IL) was used for statistical analysis. Two ways analysis of variance (ANOVA) was performed. Post hoc tests were used to determine where statistically significant differences were located among the groups (Bonferroni, Tukey). The level of significance, unless otherwise indicated, was P≤0.01.
Results

_Stathmin is expressed by migrating GnRH neurons in situ:_

Stathmin expression in the GnRH system was studied in mouse at embryonic stages E12 and E18 and in young adults (P30) by immunohistochemistry. The analysis was performed using either single-labelling immunohistochemistry on consecutive sections (8 µm apart) or double-labelling immunofluorescence on the same section. We showed that GnRH neurons migrating through the nasal mesenchyme during early stages of development express stathmin (Fig. 1).

At E12, stathmin-immunoreactivity (-ir) was detected in cells emerging from the presumptive vomeronasal organ and migrating in cords-like clusters toward the forebrain (Fig. 1A). To assess the relationship between GnRH and stathmin immunolabelling, the consecutive section was stained with anti-GnRH antiserum (Fig. 1B).

Stathmin staining pattern paralleled the GnRH neuronal distribution in the nose at this developmental stage. GnRH-ir overlapped partially with that for stathmin even though stathmin expression appeared not to be restricted to GnRH neurons. Confocal analysis of double labelling reaction on sagittal brain sections at the same embryonic stage confirmed the colocalization of GnRH and stathmin in the same cell body (Fig. 1C-E). At E18 such colocalization was still detectable in GnRH neurons crossing the crybriform plate, entering the developing forebrain and moving through the basal forebrain directed toward the hypothalamic regions (data not shown). At this stage, within these latter locations most GnRH neurons are stathmin-negative (Fig. 1F-H). Accordingly, at P30, when all the GnRH neurons are fully differentiated, no stathmin-ir can be detected in GnRH cells in any brain regions (data not shown). These results show that GnRH neurons express stathmin and that this expression is restricted to their migratory process being downregulated as the
cells reach their final destinations. To further assess the relationship between stathmin expression and the migratory activity of GnRH neurons we switched to an *in vitro* model using the GnRH immortalized cell line, GN-11.

**GN-11 cells express stathmin:**

In order to use the GN-11 cell line for functional studies we verified that this system retained expression of stathmin comparable to that observed *in vivo*.

RT-PCR (Fig. 2A), western-blots (Fig. 2B) and immunocytochemistry (not shown), demonstrated that stathmin is constitutively expressed by GN-11 cell line. RT-PCR reactions for stathmin transcripts yielded predicted 612 bp amplicons (Fig. 2A) and western blotting analysis evidenced the expression of the 19 kDa stathmin protein (Fig. 2B) in GN-11 extracts. Olfactory bulb extracts were used as positive controls (28) and GAPDH amplification confirmed that equal amounts of RNA were included in each RT-PCR reaction.

**Stathmin transfections on GN-11 neurons modify the level of stathmin expression:**

The cDNA coding for stathmin (produced by RT-PCR) was cloned in pGEM-T vector and sequenced. The amplified cDNA differs from the published sequence (J04979) for nucleotides 210 and 526, but encodes for stathmin protein with 100% amino acidic sequence homology. The amplified cDNA was subcloned in sense or antisense direction in pIRESpuro2 expression vector and transfected in GN-11 cells. Twenty-three sense (STMN+), seven antisense (STMN-) and twenty-five empty vector (pIRES) stable clones were selected by puromycin treatment and analysed for stathmin expression at RNA (Fig. 3A) and protein level (Fig. 3B). Transfection of the vector containing sense stathmin cDNA allowed transcription of stathmin mRNA, leading to an increase in stathmin protein content (STMN+) compared to control (pIRES), whereas transfection of the vector containing
antisense stathmin cDNA, leading to production of a mRNA complementary to the endogenous mRNA, knocked-down significantly stathmin expression in these transfected cells (STMN-; Fig. 3B). To exclude unspecific effects of the stathmin expression vectors, STMN transfectants were analysed the for transcript levels of stathmin-unrelated genes, such as the two tyrosine-receptors c-Met (38) and ErbB2 (39, 40) and the transcription factor AP-2α (41), which are expressed in the GN-11 parental cells. As shown in Fig. 8B, c-Met transcript levels were unchanged in between STMN-, STMN+ and pIRES cells. Unaltered levels were also observed for ErbB2 and AP-2α (not shown).

*Stathmin down-regulation modifies cell proliferation*

The proliferation rate of stathmin transfectants was analysed using crystal violet staining (see material and methods). Clones in which stathmin was down-regulated (STMN-) exhibited a proliferation activity significantly reduced as compared to control (pIRES; Fig. 4). Conversely, no significant difference in the growth rate of STMN+ clones was observed as compared to control. The growth rate of control (pIRES) was similar to that of untransfected parental GN-11 cells (not shown).

*Stathmin expression and cell morphology*

Transfection of stathmin constructs induced significant morphological changes. STMN+ cells displayed a spindle-shaped body (bipolar) with the extension of two leading processes, whereas STMN- displayed a multipolar morphology (Fig. 5A,E). pIRES cells, which express an intermediate level of stathmin compared to STMN+ and STMN- clones, exhibited both bipolar and multipolar morphologies (Fig. 5C). Quantitative morphological analysis was performed, taking into account diameter, area and perimeter of random cells (n=249 for STMN+ cells; n= 228 for pIRES cells; n=323 for STMN- cells) for each clone (Fig. 6). The mean ratio between the maximum and minimum diameter (± SEM) was
calculated. These values were 4.8 ± 0.14 µm for STMN+ clones, 3.2 ± 0.11 µm for pIRES and 2.6 ± 0.05 µm for STMN-, confirming that a significant difference in cell morphology existed between the three clones of transfected cells (P<0.01).

Mean cell perimeters (± SEM) of clones STMN+, pIRES and STMN- were 171.3 ± 3.1 µm; 155.9 ± 2.4 µm; 147.8 ± 3.2 µm respectively. Statistical analysis evidenced a significant difference (P<0.05) of cell perimeters between STMN+ and pIRES, STMN- and pIRES, STMN+ and STMN- clones respectively. Mean cell body areas (± SEM) of clones STMN+, pIRES and STMN- were 798.8 ± 41.6 µm; 857.8 ± 25.4 µm; 987.2 ± 54.7 µm. Statistical analysis showed significant difference (P<0.05) between STMN+ and STMN- clones.

To test whether the overexpression of stathmin is associated with abnormal MTs, STMN cells were immunolabelled for α-tubulin. We observed different organizations of this cytoskeletal component between clones, which could account for STMN transfectants shape; however, the density of microtubules was not modified (Fig.5 B,D,F).

Stathmin level affects two dimensional motility

Being interested in studying stathmin-mediated migration, we performed a first round of experiments to determine whether differences of stathmin expression could modulate motility, here defined as two dimensional movement, or chemokinesis. The nuclear coordinates of single cells were measured every 30 minutes for 3 hours and the mean speed of cell movement was calculated. Our results show different chemokinetic responses: stathmin over-expression induced a significant increase in motility (23.6 µm/h) when compared to the pIRES (13.5 µm/h), whereas no significant difference between STMN- (12.6 µm/h) and pIRES cells motility was detected.
Stathmin level modifies aggregation ability

Parental GN-11, pIRES and STMN- cells, in subconfluent culture conditions, exhibit cell-cell adhesion, while STMN+ cells grow exclusively as single cells. Using the *hanging drop* approach (35) we showed that, unlike pIRES and STMN-, STMN+ cells were scattered and spread into the collagen gel and do not aggregate at all (Fig. 7A). To further investigate a possible role for stathmin in the impairment of cell-cell adhesion, we analysed N-cadherin protein expression in the different clones. As shown in Fig. 7B, we found that STMN- cells expressed higher levels of full length N-cadherin (130 kDa) as compared to pIRES cells and a truncated (95 kDa) isoform (42) which was absent in the STMN+ and pIRES clones. Concurrently with the above observations, STMN+ cells express very low level of full length N-cadherin (130 kDa).

Stathmin level modifies three dimensional basal and chemotactic motility

To evaluate the ability of STMN transfectants to migrate in a three dimensional environment, we assayed basal motility and chemotaxis using transwell assay, a common approach for sensitive measurement of cellular response to specific chemotropic signals (43; 44). In previous studies, we and others have shown that GN-11 cells display migratory activity (36; 45). To determine whether stathmin expression was involved in GN-11 basal cell migration, transwell assays were performed with STMN+ and STMN- clones in serum free medium. The statistical analysis of migrated cells after 6 hours of incubation clearly showed the different migratory activity of pIRES, STMN+ and STMN- clones (Fig. 8). STMN+ cells exhibited a significant increase in cell migration activity when compared to pIRES, whereas, in the same experimental conditions, STMN- cells showed a significant decrease.
STMN clones were then assayed for their ability to respond to the chemotactic molecule, hepatocyte growth factor/scatter factor (HGF/SF). We have previously shown that GN-11 cells express the tyrosine kinase receptor c-Met and respond to the chemotactic stimulus induced by its ligand, HGF/SF (45). Cells were seeded in the transwell and exposed to a source of HGF/SF (50 ng/ml; 45) which was added in the bottom chamber (see materials and methods). As shown in Fig. 8A, all clones responded significantly to HGF/SF treatment, with an increase in cell migration as compared to SFM treatment. Noteworthy, the amplitude of the response is correlated to the basal motility of each clone. In addition, STMN+ cells displayed increased cell motility both in SFM condition and in the presence of HGF/SF when compared to pIRES cells. Concurrently, STMN- cells displayed decreased cell motility both in SFM condition and in the presence of HGF/SF when compared to pIRES cells. Finally, no significant differences in c-Met expression were observed between GN-11, pIRES, STMN- and STMN+ cells, as detected by RT-PCR and immunoblotting analyses (Fig. 8B).
Discussion

In the present paper, we show that the microtubule (MT) associated phosphoprotein stathmin is expressed in vivo in migrating GnRH producing cells during development as well as in vitro, in the GN-11 cells (30). In addition, we demonstrate that the modulation of stathmin expression significantly affects shape and motility of GN-11 cells.

During CNS development and in the adult, MTs are involved in numerous intracellular events, such as proliferation, differentiation and migration. In neurons, MTs play specific roles in axonal outgrowth, pathfinding, and synapse formation, as well as in axonal and dendritic transport (46). We have focused our interest on the MT-associated protein stathmin and investigated its role as one of the possible regulators of these events.

Stathmin phosphoprotein interferes with MT assembly and plays an important role in the regulation of MT dynamics during cell cycle progression. It is also highly expressed in early embryos (47; 48; 49), gonads (50) and in particular it has been shown to be present in several brain areas (28; 51; 52; 53; 54), but its function in the CNS is not completely clear. Stathmin has also been proposed to act as an intracellular relay for extracellular signals (55) and many proteins, apart from tubulin, have been identified as target/partners for stathmin (56; 57; 58). A stathmin gene has also been identified in Drosophila (59); like the mammalian protein, Drosophila stathmin interferes with MT dynamics both in vitro and in vivo. Furthermore, functional inactivation of the stathmin gene in Drosophila, by RNA interference, leads to abnormal germ cell migration in the embryo and to dramatic defects in the formation of the nervous system, supporting a direct role of stathmin in these essential biological processes.

In mammals, stathmin immunostaining in the CNS localizes to immature olfactory neurons as well as to migrating cells generated from the olfactory epithelium, supporting
the role of this protein in neurogenesis and cell migration (29). Although stathmin had been associated with numerous cell events, its biological role remained elusive as inactivation of the stathmin gene in mouse resulted in no clear deleterious phenotype (60). In a recent study, microinjections of stathmin antisense oligonucleotides in the lateral ventricle of adult rats, inhibited interneuronal migration from the SVZ to the olfactory bulb via RMS (61).

We now evidence, by consecutive and simultaneous double immunohistochemical analyses, that stathmin is expressed by GnRH neurons. This expression is specific of the migration step and ceases as soon as GnRH neurons reach their final hypothalamic destination. Thus, stathmin expression in GnRH neurons is temporally and spatially regulated during mouse development. Based on the pre-existing literature and on the present results, we postulated that stathmin is involved in the control of migration properties of GnRH neurons. To determine whether a correlation might exist between stathmin expression level and migratory capabilities of GnRH neurons, we moved to GN-11 cell line. This cell line was derived from a tumour developed in the cribriform plate of mouse embryos (30). GN-11 cells retain many of the morphological and behavioural features of migrating neurons (59; 45; 63). Therefore, this cell line provides a suitable model to study the involvement of stathmin in the regulation of GnRH neuronal migration.

Complete stathmin cDNA was cloned both in sense and antisense orientation in pIRES vector and stably transfected into GN-11 cells. Selected clones showed over-expression (STMN+) or down-regulation (STMN-) of stathmin as compared to control clones. As already demonstrated for K562 erythroleukemia cells (23) and for human embryonic kidney 293 cells (64), the alteration of stathmin expression in GN-11 cells induces modifications in proliferation activity, confirming a role for stathmin as mitotic regulator in this cell line. Interestingly, beside its effect on proliferation, modifications of stathmin expression caused also changes in GN-11 cell morphology. High levels of stathmin expression induced the cells to adopt the typical morphology of in vivo GnRH
migrating neurons (65), with elongated and spindle-shaped cell bodies and the extension of a leading and a trailing process. Cells expressing low levels of stathmin showed neurite-like outgrowth and displayed a multipolar morphology. The significance of these morphological observations was evaluated and confirmed by quantitative analysis of parameters such as area, perimeter and shape.

Bidimensional analysis of cell motility revealed a significant increase in the speed of STMN+ cells when compared to control and ST MN- cells. Interestingly, STMN+ cells did not establish any contact with each other, suggesting a possible variation of the expression of cell-cell adhesion molecules. Previous studies have shown that cell adhesion molecules, including N-cadherins, are implicated in the formation of the nervous system (66). Their expression is highly regulated during nervous system development to control cell migration, neurite outgrowth, fasciculation, and synaptogenesis. It has also been demonstrated that down-regulation of cadherin’s expression hastened the migration of newly generated neurons produced in the subependymal zone (67). In other model systems, inactivation of the cadherin adhesion complex seems to be associated with cell dedifferentiation, invasion and regional metastasis (68; 69), all processes requiring cell motility. Based on these data, we focused our attention on the adhesive properties of GN-11 cells and analysed the expression of N-cadherins in our experimental model. Interestingly, we found out that the expression patterns of stathmin and N-cadherins are inversely correlated: overexpression of stathmin induces a down-regulation of N-cadherin expression and the loss of the ability to form cell aggregates by the hanging-drop technique, whereas down-regulation of stathmin is accompanied by an increase in N-cadherin protein expression and cell-cell adhesion. A correlation between stathmin and cadherins has been previously highlighted by Balogh and co-workers, who showed that cell-cell contacts, probably mediated by cadherins, may be important in the control of stathmin expression (70).
Using transwell assay, we showed that spontaneous migratory activity of GN-11 cells is sensitive to stathmin content. Indeed, transfection of GN-11 cells with stathmin sense construct significantly increases basal migration, whereas down-regulation of stathmin expression by antisense construct transfection inhibits such activity when compared to control cells. These results are in accordance with those of Jin and co-workers demonstrating a role for stathmin in the RMS neuronal migration by ventricular antisense oligonucleotide injection (61).

HGF/SF motogenic activity has been demonstrated in several cell lines, during embryogenesis and in metastatic tissues (for a review see 38). In the developing CNS, HGF/SF and its receptor c-Met are involved in the migration of cortical (71) and GnRH neurons (45). In addition, GN-11 cells have been previously shown to express c-Met and display chemotactic responsiveness to HGF/SF stimulation (45). Since the level of c-Met expression is not affected by stathmin content, the migratory activity of STMN+, pIRES and STMN- transfectants has been measured under the exposure to HGF/SF. A chemotactic response to HGF/SF is observed for all transfectants and the amplitude of the response correlates to stathmin content. The activation of c-Met receptor by HGF/SF leads to the recruitment, among others, of the scaffolding protein Grb2 which activates Rac-1 (38). On the other hand, the activation of Rac-1 dependent kinase has been recently demonstrated to regulate stathmin activity at the leading edge of migrating cells (72; 73). Therefore, a challenging hypothesis that will deserve further investigation, is the regulation of stathmin activity by Rac-1 dependent kinase in GN-11 cells in response to HGF/SF.

The temporal expression of stathmin protein by migrating GnRH neurons is in accordance with the in vitro findings that we obtained using GN-11 cell line. In addition, once GnRH neurons have reached their final hypothalamic destinations, they no longer express stathmin and complete their differentiation to become integral components of the hypothalamic-pituitary-gonadal axis, which is essential for establishment of reproductive
competence. Whether down-regulation of stathmin expression is a necessary step for GnRH neuronal differentiation or, inversely, a consequence of the differentiation program of the neuroendocrine cells, deserves further investigations.
References


8. Ronnekleiv OK and Resko JA. 1990 Ontogeny of gonadotropin-releasing hormone-containing neurons in early foetal development of rhesus macaques. Endocrinology, Jan; 126, 498–511


43. Isenberg JS. 2003 Inhibition of nitric oxide synthase (NOS) conversion of L-arginine to nitric oxide (NO) decreases low density mononuclear cell (LD MNC) trans-endothelial migration and cytokine output. J Surg Res. Sep;114(1):100-6


71. Powell EM, Mars WM, Levitt P. 2001 Hepatocyte growth factor/scatter factor is a motogen for interneurons migrating from the ventral to dorsal telencephalon. Neuron, Apr 30(1):79-89


**Figure legends:**

**Fig.1:** Stathmin and GnRH in vivo immunohistochemistry.
Panel A and B: immunohistochemistry for stathmin (A) and GnRH (B) performed on consecutive sagittal sections of a E12 mouse embryo. Immunoreactive cells emerged from the developing olfactory epithelium and migrate in chains through the olfactory mesenchyme toward the forebrain. Stathmin immunoreactivity resembles the staining pattern of GnRH. Note that the stathmin antibody likely labels the same groups of migrating elements stained with the GnRH antiserum. Abbreviations: *fb*, forebrain; *n*, nose; *op*, olfactory placode; *t*, tongue. *Scale bars: * 500µm, ** 100µm.*
Panel C-E: double simultaneous immunohistochemical staining for stathmin (C) and GnRH (D) at E12 in the nasal mesenchyme. A complete colocalization between stathmin and GnRH is shown in panel E (*merge* of C and D). *Scale bars: 10µm.*
Panel F-H: double simultaneous immunohistochemical staining for stathmin (F) and GnRH (G) at E18 in the hypothalamus. As shown in panel H (*merge* of F and G), no colocalization between stathmin and GnRH was observed in this region. *Scale bars: 10µm.*

**Fig.2:** Stathmin expression in GN-11 cell line extracts. **Panel A:** Total GN-11 mRNA was assayed by RT-PCR for stathmin, GAPDH served as control. Predicted amplicons molecular weights were: 612 bp for stathmin and 188 bp for GAPDH. **Panel B:** GN-11 cells protein extracts were analysed for stathmin (19 kDa) expression. Abbreviations: *MW*, molecular weight; *OB*, olfactory bulb (positive control).

**Fig.3:** Representative stathmin expression in STMN stable transfectants. **Panel A:** Total mRNAs were assayed by RT-PCR for stathmin, GAPDH served as control. Predicted amplicons molecular weights were: 612 bp for stathmin and 188 bp for GAPDH.
Panel B: Total proteins from STMN+, pIRES and STMN- cells were assayed by western-blot analysis for stathmin. β-actin served as control of the quantity of proteins.

Fig.4: Effect of stathmin level modulation on cell proliferation in SFM culture condition. Lower levels of cell proliferation were found in conditions of lower level of stathmin (STMN- cells).

Abbreviation: O.D., optical density (see materials and methods section).

Fig.5: Morphological characteristics of STMN stable transfectants. STMN+ cells displayed a spindle-shaped body with the extension of two leading processes (A), while STMN- were much more spread (E). pIRES cells (C), which express an intermediate level of stathmin compared to STMN+ and STMN- clones, exhibited both spindle (white arrow) and spread (black arrow) morphology. All the STMN transfectants (B,D,F) exhibit intact microtubule networks, as shown by α-tubulin-ir. Scale bars: 20µm

Fig.6: Quantitative analyses of STMN stable transfectants morphological parameters. The morphometric data of every cell soma were measured semi-automatically: cell body boundaries were manually traced. For each cell body, spindle-shaped morphology (A)(evaluated as ratio between maximum and minimum diameter), perimeter (B) and area (C) were analysed. Each value represents the mean ± standard error; n= 249 STMN+ cells; n=228 pIRES cells; n=323 STMN- cells. Statistically significant differences of each group with one another were observed when comparing data concerning spindle-shaped morphology and perimeter (A, B). When comparing data concerning areas, statistically significant differences were observed only between STMN+ and STMN- cells (C).
**Fig.7:** Panel A: Aggregation assays of STMN+, pIRES and STMN- cells. pIRES and STMN- cells form aggregates in collagen gel (*hanging drop technique*), while STMN+ cells are scattered and spread into the collagen gel. The inset shows, at higher magnification, STMN+ cells not forming aggregates.

Panel B: N-cadherin protein expression in STMN clones. STMN- cells, compared to pIRES, express high level of full length N-cadherin (130 kDa) and, in addition, the induction of a truncated (95 kDa) isoform. Concurrently, STMN+ cells express very low level of full length N-cadherin (130 kDa).

**Fig.8:** Panel A: Three dimensional migration assay. Chemotactic response of STMN+, pIRES and STMN- cells, performed using the *transwell method*: $10^5$ cells were allowed to migrate for 6 hours in the absence or in the presence of 50 ng/ml HGF/SF. Each value represents the mean $\pm$ standard deviation of three experiments in triplicate. All clones respond significantly to HGF/SF treatment, with an increase in cell migration when compared to untreated conditions and the amplitude of the response is correlated to the basal motility of each clone. STMN+ cells, when compared to pIRES cells, display an increased motility both in SFM condition and in the presence of HGF/SF. STMN- cells, when compared to pIRES cells, display a decreased motility both in SFM condition and in the presence of HGF/SF.

Panel B: c-Met expression in STMN transfectants extracts. Total mRNAs were assayed by RT-PCR for c-Met, GAPDH served as control. Predicted amplicons molecular weights were: 519 bp for c-Met and 188 bp for GAPDH. Protein extracts were analysed for c-Met (145 kDa) expression. Abbreviations: *MW*, molecular weight; *SFM*, serum free medium.
(A) MW  STMN+  pIRES  STMN-

STATATHMIN
612 bp

GAPDH
188 bp

(B) STMN+  pIRES  STMN-

STATATHMIN
19 kDa

β-ACTIN
42 kDa
(A) SPINDLE-SHAPED MORPHOLOGY

(B) PERIMETER

(C) AREA