Research Article

Ret Finger Protein: An E3 Ubiquitin Ligase Juxtaposed to the XY Body in Meiosis

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During prophase I of male meiosis, the sex chromosomes form a compact structure called XY body that associates with the nuclear membrane of pachytene spermatocytes. Ret Finger Protein is a transcriptional repressor, able to interact with both nuclear matrix-associated proteins and double-stranded DNA. We report the precise and unique localization of Ret Finger Protein in pachytene spermatocytes, in which Ret Finger Protein takes place of lamin B1, between the XY body and the inner nuclear membrane. This localization of Ret Finger Protein does not seem to be associated with O-glycosylation or sumoylation. In addition, we demonstrate that Ret Finger Protein contains an E3 ubiquitin ligase activity. These observations lead to an attractive hypothesis in which Ret Finger Protein would be involved in the positioning and the attachment of XY body to the nuclear lamina of pachytene spermatocytes.

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1. Introduction

During meiosis, homologous recombination between chromosomes is allowed by synopsis of chromosome’s pairs during pachytene stage. At this stage, all the chromosomes are attached to the nuclear membrane by their telomeres. In order to deal with asynchronous synopsis of the XY chromosomes, mammals have adopted a strategy that involves chromatin condensation and seclusion of the sex chromosomes into a specialized and visibly distinct nuclear territory, known as the XY body [1, 2]. After early pachytene, the XY chromosomal pair becomes highly condensed, transcriptionally silent, and positioned adjacent to the nuclear envelope.

Localization of the condensed XY body to one part of the nucleus suggests that a particular composition of the nuclear matrix could be involved in the positioning and/or attachment of the XY body or sex chromosomes to the nuclear lamina [3, 4]. The nuclear lamina provides an attachment site for chromatin and facilitates processes associated with DNA replication and transcription. In germ cells, lamin B1, as well as specific isoforms of lamin A/C, is evidenced in the nuclear matrix [5, 6].

The precise molecular events underlying sex chromatin condensation and meiotic inactivation of sex chromosomes remain to be elucidated. However some hypotheses have been advanced based on the presence or the absence of specific proteins inside the XY body [2]. Matsuura et al. [7] reported that SUMO-I and PIAS-y are found inside the XY body and proposed that Ret Finger Protein (RFP) can also be associated with the XY body and could then have a role in the silencing of sex chromosomes.

The initial interest in RFP arose from the demonstration that it becomes oncogenic when fused with the RET tyrosine kinase [8]. In healthy animals, RFP is highly expressed in male germ cells, in particular in primary spermatocytes and in round spermatids [9]. RFP is usually detected as a 58 kDa protein by western blot analysis. However, two isoforms of RFP, 58 kDa and 68 kDa, can readily be detected in testis [9, 10]. Tezel et al. [11] have shown that the 68 kDa isoform results from an O-glycosylation of RFP. Despite extensive studies, the biological functions of RFP remain unclear. It has been described as a nonspecific DNA-binding protein strongly associated with the nuclear matrix [10]. RFP has also been shown to partially colocalize with PML and Inf6 [12, 13]. RFP has been implicated in general...
transcriptional repression mechanisms, in particular when it interacts with the Enhancer of Polycomb (EPC), Mi-2β and histone deacetylase 1 (HDAC1) [14–16]. In somatic cells, SUMO-I modification of RFP induces its relocalization into nuclear bodies reminiscent of the PML bodies and strengthens its transcriptional repressor activity [7]. RFP has also been proposed to repress transcriptional activity of specific proteins such as Retinoblastoma protein or bHLH transcription factors [17, 18]. However, the role of RFP in meiotic cells remains unclear. Finally, RFP belongs to the family of RING-B-Box-Coiled Coil-B30.2 proteins [19]. According to the International Mouse Genome Informatics Committee, the proteins containing RING-B-Box-Coiled Coil motif have been renamed TRIM proteins, and the name of TRIM 27 was assigned to RFP [20, 21]. As a result of numerous studies, it has been proposed that all the RING finger containing proteins, as well as TRIM proteins, could act as E3 ubiquitin ligases and be involved in proteasomal degradation process (reviewed in [20, 22, 23]).

We report here the precise localization of RFP in pachytene spermatocytes. We demonstrate that RFP exhibits a unique and specific localization that may define the positioning of the XY body. Moreover, as predicted by the presence of a RING domain, we demonstrate that RFP possesses an E3 ubiquitin ligase activity.

2. Materials and Methods

2.1. Cells and Virus Propagation. 15P-1 cells [24] were maintained at 32°C in DMEM media (Gibco: 41966-029) supplemented with 10% Fetal Bovine Serum (FBS, Gibco: 10270-098) and 1/100 Penicillin/Streptomycin (P-S, Gibco: 15140-163).

S9 cells (BD Biosciences: 554762) were maintained in TNM-FH media supplemented with 1/100 P-S as previously described [25]. Baculoviruses were propagated, prepared, and tittered in S9 cells as described by the manufacturer (BD Biosciences).

Fractionation of germ cells from twenty B6D2F1 mice by elutriation centrifugation was performed as previously described [26] and the purity of each fraction was determined by elutriation centrifugation was performed as previously described [26] and the purity of each fraction was determined by DNA staining with DAPI. Fractions used were estimated to contain more than 80% of purified spermatocytes, round spermatids, or elongated spermatids.

2.2. Construction of Plasmids and Baculoviruses. Primers RFPsense139 5′ CCAGAACTCATGCTGCAGCCAGCAG and RFPrev569 5′ CCGCTAGGACTTGGTCTGCCG-GTGCCT were used to amplify the RING-B-Box domain of RFP from total testis cDNA. EcoRI/XhoI digested PCR fragment was ligated in pGex4T-1 (Amersham Biosciences). RFP coding sequence (CMV) and an additional Myc tag.

2.3. Protein Purification in Bacterial Cell System. BL21 bacterial cells transformed with pGEX vectors were amplified at 37°C in LB medium supplemented with 0.1 mM Zn²⁺ and 50 μg/mL ampicillin. At optical density 0.6–0.8, cells were induced at 30°C by adding 1 mM IPTG for two hours. GST fusion proteins were purified using Glutathione Sepharose 4B (Amersham Biosciences: 17-0756-01), eluted with reduced glutathione (Sigma: G-4251), and dialyzed against PBS (Phosphate-Buffered Saline pH 7.4) containing 1 μM ZnSO₄ using disposable dialyzer (Sigma-Aldrich: Z368385-10EA).

2.4. TPEN Treatment of GST–RING/RFP. For cation chelation, GST–RING/RFP immobilized on Glutathione Sepharose was incubated overnight at 4°C, with 5 mM N,N,N′,N′-tetrakis(2-pyridylmethyl)-ethylenediamine (TPEN) (Sigma-Aldrich: 87641) or 1% ethanol (vehicle) in 50 mM Tris, pH 7.5. After extensive washes, one half of the immobilized GST–RING/RFP was treated with 30 μM of ZnCl₂ for 45 minutes at room temperature. After washing, samples were used for ubiquitination assays.

2.5. RFP Immunoprecipitation from Baculovirus-Infected Cells. 15P-1 cells, seeded 15 to 18 hours before infection at 80%–90% confluency, were exposed in TNM-FH media to 1 plaque-forming units per cell of wild type or Bac-Myc-RFP recombinant baculoviruses. After 2 hours, the inoculum was replaced by fresh DMEM medium containing 5 mM sodium butyrate (Sigma-Aldrich: B5887) and cells were incubated at 32°C for 72 hours. Cells were lysed in a buffer containing 50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1% NP-40, and EDTA-free “complete” protease inhibitors (Roche: 1875380) sonicated briefly, and centrifuged for 10 minutes at 13,000 rpm at 4°C (Biofuge pico, 3325 Heraeus rotor; Sorvall, Norwalk, Conn.). Supernatant was used for RFP immunoprecipitation using a rabbit polyclonal antibody to RFP (IBL Japan: JP 18791) according to the supplier’s instructions. After washing, immunoprecipitated RFP was used for ubiquitination assays.

2.6. Western Blots. Elutriated germ cells [26] were lysed. Protein concentration was evaluated using a Bradford assay (BioRad: 500-0006); 20 μg of proteins per lane were subjected
to SDS Page electrophoresis (8%) and blotted on PVDF membrane. Rabbit polyclonal antibody to RFP (IBL Japan: JP 18791) and mouse anti-rabbit peroxidase-conjugated antibody (Sigma-Aldrich: a2074) were used at 1 : 200 dilution and 1 : 10000 dilution, respectively. Immunoblots were done through enhanced chemiluminescence detection (Amersham Biosciences: RPN2132). ECL blots were analyzed using a Fuji Film Intelligent Dark box (LAS-3000) and the associated computer programs (Image Reader Las-3000 and Multi-Gauge V2.3).

2.7. In Vitro Substrate-Independent Polyubiquitination. In vitro polyubiquitination assays were performed as described in Hagglund et al. [28], in ubiquitination buffer (50 mM Tris, pH 7.5, 2.5 mM MgCl₂, 0.5 mM DTT) containing 40 ng of E1 (Calbiochem: 662070), 40 ng of one E2 (Biomol Affiniti, [His₆]-UbCH1 (uw9020); [His₆]-UbCH2 (uw9025); [His₆]-UbH3 (uw8730); [His₆]-UbCH15a (uw9050); [His₆]-UbH5b (uw9060); [His₆]-UbCH15c (uw9070); [His₆]-UbH6 (uw8710); [His₆]-UbCH7 (uw9080); [His₆]-UbH10 (uw8715)) or [His₆]-HR6A or [His₆]-HR6B both produced in the laboratory, 2 μg of biotinylated ubiquitin (Biomol Affiniti: uw8705), 0.2 mM ATP, 1 mM creatine phosphate (Calbiochem: 2380), and 15 units of porcine heart creatine phosphokinase (Calbiochem: 238395) in final volume. When indicated, the reaction was performed in absence of ATP. Immunoprecipitated RFP and either five μg of GST or GST-RING/RFP recombinant protein were added to the reaction mixture and incubated at 37°C for 90 minutes. Reactions were stopped with 4X Laemmli buffer and subjected to SDS-Page analysis. Membranes were rinsed and blocked in PBS, 0.1% BSA, and 0.1% Tween 20 and reacted 1 hour at room temperature with 0.02 unit/mL of streptavidin peroxidase (Roche: 1089153). Biotinylated ubiquitin conjugates were revealed after ECL. Immunoblots were done through enhanced chemiluminescence detection (Amersham Biosciences: RPN2132). ECL blots were analyzed using a Fuji Film Intelligent Dark box (LAS-3000) and the associated computer programs (Image Reader Las-3000 and Multi-Gauge V2.3).

2.8. Immunofluorescence Analysis. Immunofluorescence labeling has been performed on testis frozen sections, on total germ cell preparations, or on elutriated germ cells prepared as previously described [26].

Cells were fixed in cold ethanol (−20°C) for 1 hour before a 30-minute blocking step in PBS containing 1% BSA (Sigma: A-4503) and 20% normal human serum.

Colabeling experiments have all been performed in a sequential manner according to the same protocol. Antibodies were diluted in PBS containing 1% BSA and 2% normal human serum. After each antibody incubation, samples were washed three times with PBS containing 1% BSA. Primary specific antibodies were incubated overnight at 4°C and the secondary antibody was added for 1 hour at room temperature. The second specific antibody was then incubated for 2 hours at room temperature and the secondary antibody was added for 1 hour. After a final wash in PBS, slides were mounted in Vectashield reagent with DAPI (Vector Laboratories, Inc., Burlingame).

The specific antibodies were: a goat polyclonal IgG Lamin B1 antibody (Santa Cruz Biotechnology: M-20, sc-6217); a rabbit polyclonal RFP (IBL Japan: JP 18791) antibody; a mouse monoclonal SUMO-I antibody (Zymed laboratories: 33-2400), a mouse IgG anti phosphohistone H2AX antibody (Stress gen Bioreagents corp., Canada: KAM-CC255) respectively diluted at 1 : 80, 1 : 150; 1 : 400 and 1 : 800.

The secondary antibodies (dilution 1 : 400) were: a Cy-3 conjugated affinity Pure donkey anti-goat IgG (Jackson Laboratories: 705-165-147); a goat anti-rabbit Alexa 516 and a goat anti-mouse Alexa 546 antibodies (Molecular Probes, Oregon: A11035 and A11030).

2.9. Image Analysis. Image acquisition (Objective 60 × 1.4 Uplanapo) and deconvolution using the constrained iterative algorithm (Softworx 2.5) was performed with Applied Precision DeltaVision system (Applied Precision, Issaquah, WA) built on an Olympus IX-70 base.

In Figures 4(a) and 4(b) images were made by the sum of the intensities of three consecutive images taken in Z position (0.15 nm interval) in each channel. Treatment was performed using ImageJ NIH software. The 3D reconstruction (Figures 4(c), 4(d), and 4(e)) has been made with 30 consecutive images taken in Z position and the reconstruction is made using Velocity software and Image plugin Volume Viewer (Kai Uwe Barthel).

2.10. Immunogold Analysis. Samples fixation, ultrathin sections and immunogold labeling was performed as previously described [29]. Anti-RFP and anti-SUMO-I antibodies were diluted 1 : 100 in PBS; 1% BSA; 1% NGS. Colloidal gold conjugated anti-rabbit IgG antibodies (15 nm) and colloidal gold conjugated anti-mouse IgG antibodies (10 nm) were diluted 1/20 in PBS 0.01% fish-skin gelatin. Preparations were observed with a Phillips CM12 electron microscope operating at 80 kV. No signal was observed when the first antibody was omitted.

3. Results

3.1. RFP Exhibits Various Localizations in Meiotic and Postmeiotic Germs Cells. RFP localization in mouse male germ cells, and in particular in pachytene spermatocytes, has previously been described as a perinuclear cap structure [9]. Immunolabeling of RFP in testis sections corresponding to stages late VIII and IX of mouse spermatogenesis (Figures 1(a)–1(c)) revealed at least two different localizations for RFP in germ cells: a thin cap-like structure forms next to the nucleus of meiotic germ cells which then seems to condense to a spot in later stages of spermatogenesis. Optical and electronic analysis indicate that these foci of RFP are then found in residual bodies (data not shown).

Two isoforms of RFP 58 kDa and 68 kDa have previously been described in testis [9]. Tezel et al. [11] have shown that the 68 kDa form resulted from an O-glycosylation of RFP. Considering the specific localization of RFP protein in the nuclear matrix of pachytene spermatocyte, we checked
the possible relationship between this posttranslational modification and the localization of RFP. We investigated the respective distribution of the 58 kDa and 68 kDa isoforms in elutriated germ cell populations (Figure 1(d)). Our results indicated that maximal RFP expression is found as expected in round spermatids [9]. However, both isoforms were found in pachytene spermatocytes, in round spermatids, and in elongated spermatids. Furthermore, the ratios between the two isoforms were equivalent in these different germ cell populations (53.43%/46.57% in pachytene spermatocyte, 51.68%/48.32% in round spermatids and 49.88%/50.12% in elongated spermatids for the ratio of 58 kDa/68 kDa isoform, resp.). Relative distribution of each RFP isoform does not show any clear relationship between the level of O-glycosylation and the localization of RFP in pachytene cells.

3.2. RFP Defines a Region Underlying the XY Body in Pachytene Spermatocyte. It has been reported that PIASy and SUMO-I localizes within the XY body [7]. In order to examine the precise localization of endogenous RFP compared to XY body markers, such as SUMO-I and γH2AX proteins [30, 31], we have performed colabeling experiments at the structural and the ultrastructural levels on meiotic spermatocytes. We demonstrate that RFP localized as a small cap at the periphery of the pachytene spermatocyte nuclei (Figure 2). Colabeling experiments made (Figures 2(d) and 2(e)) indicated that RFP and γH2AX proteins did not
colocalize but rather defined two different domains around and within the XY body. Specifically, γH2AX is detected inside the XY body and RFP is a distinct layer adjacent to the XY body, on the side facing the nuclear membrane. These results were confirmed using anti-RFP and anti-SUMO-I antibodies giving a similar pattern (Figures 2(f) and 2(g)). To our knowledge, RFP is the only protein that has been shown to exhibit such a particular localization.

Electron microscopy analysis of immunogold labeling, performed on testis sections using anti-RFP and anti-SUMO-I antibodies, confirmed that the two proteins were very closely apposed but did not colocalize within spermatocyte nuclei (Figure 3). In order to obtain semiquantitative data, we have evaluated the number of 15 nm gold beads (corresponding to RFP) and the number of 10 nm gold beads (corresponding to SUMO) in Figure 3. To compare the localization of SUMO and RFP, we have evaluated the number of gold beads located in the nucleus and in the XY body. In Figure 3(a), nuclear surface has been evaluated to 59.48 μm² and the surface of the XY body to 2.68 μm². By counting the beads in the nucleus, we found a concentration of 0.25 beads/μm² for SUMO labeling and of 0.17 beads/μm² for RFP labeling. Within the XY body, we have obtained 9.32 beads/μm² for SUMO labeling and 0 beads/μm² for RFP labeling. Comparable results were obtained when Figure 3(b) was analyzed in the same way. Moreover, in Figure 3(a), the total length of nuclear envelope has been evaluated to 23.40 μm and the membrane facing the XY body to 2.75 μm. Three 15 nm gold beads have been found localized at the nuclear membrane facing the XY body. So, these observations confirm that RFP is localized at the border of the nucleus facing the XY body, while SUMO-I is inside the nucleus, associated with the XY body. These results were summarized in Figure 3(e).

### 3.3. RFP Takes Place of Lamin B1 in Region of the Nuclear Membrane Adjacent to the XY Body

We have noticed that the labeling of the nuclear membrane of pachytene spermatocyte with an anti-lamin B1 antibody is interrupted in a region reminiscent of the size and the position of RFP and XY body [5]. Colabeling experiments using total germ cell preparation, anti-lamin B1 and anti-RFP antibodies showed that RFP accumulated in a region where lamin B1 was almost completely absent (Figures 4(a) and 4(b)). After deconvolution and 3D reconstruction, we visualized RFP labeling as a disc inside the nuclear matrix (Figures 4(c)–4(e)). When colabeling experiments were done with lamin B1 antibody and either SUMO-I or γH2AX antibodies, we could confirm that this region of the nuclear envelope adjacent to the XY body was devoid of lamin B1 labeling (data not show). According to our results, we conclude that in the nuclear lamina of pachytene spermatocyte, the absence of lamin B1 could be involved in the specific localization of RFP adjacent to the XY body, and thus, RFP could provide a specific attachment point for sex chromosomes.

### 3.4. RFP Has an E3 Ubiquitin Ligase Activity Associated with Its RING Domain

In somatic cells, RFP has been involved in transcriptional regulation. However, regarding its main localization in meiotic and postmeiotic germ cells, we looked for another function for RFP. RFP contains a RING domain and RING finger proteins have been proposed to be involved in degradation via the proteasome [28, 32, 33].
Figure 4: Relationship between XY body, RFP, and nuclear matrix in pachytene spermatocyte. Coimmunofluorescent labeling of lamin B1 (green) with RFP (red). (a), (b) At this cell stage, RFP takes place of lamin B1 along the border of the nucleus (DAPI, blue; bar, 8 μm). (c)–(e) Data presented as volumes within a 3D-(xyz)-space reconstructed from an RGB image stack. The three dimensions are represented by oriented arrows (x: yellow, y: white, and z: red arrows), considering xyz reference positions taken in (c).

We first tested the ability of an Myc-tagged RFP to promote an in vitro substrate-independent polyubiquitination [28]. As shown in Figure 5(a), a dark smear, characteristic of polyubiquitination, can be observed with specific E2 enzymes such as UbcH1, UbcH5A, 5B, 5C, UbcH6, and HR6B.

Using a GST-RING/RFP fusion protein, we demonstrate that the RING domain isolated from RFP contained the E3 ubiquitin ligase activity. This activity was strictly dependent on ATP hydrolysis (Figure 5(b)). The mandatory presence of Zn2+ was demonstrated by using TPEN, a chelator of divalent cations, which abolished the E3 activity (Figure 5(c), left panel). As expected, after readdition of an excess of Zn2+, the GST-RING/RFP retrieved its polyubiquitination activity (Figure 5(c), right panel). Control treatments with vehicle only or vehicle plus Zn2+ did not affect the E3 activity of the GST-RING/RFP protein (data not shown).

4. Discussion

We report here that in pachytene spermatocytes, RFP defines a precise and restricted region of nuclear matrix that correlates with the position of the XY body. This observation is supported by colabeling experiments using both deconvolution and electron microscopy. To our knowledge, the localization of RFP, associated with the nuclear matrix in a region juxtaposed to the XY body but not within, is unique in regard to the XY body. Of the lamins expressed in somatic cells, lamin B1 is the only one that could be detected in spermatocytes [5, 6]. Colabeling experiments using anti-lamin B1 and anti-RFP antibodies indicate that RFP occupied a region in the nuclear matrix, facing the XY body and devoided of lamin B1. Regarding the ability of RFP to interact with double-strand DNA and with nuclear matrix proteins [10], our data suggest that RFP could be involved in both positioning of the XY body in pachytene matrix spermatocyte and attachment of this structure to a region of the nuclear envelope matrix devoid of lamin B1. Whether this is mediated through strong association of RFP with other nuclear lamins or other proteins present in the XY body remains to be determined. However, it is unknown whether the localization of RFP is causative of XY body localization or is the result of the apposition of the XY body close to the nuclear membrane. Moreover, one cannot exclude that RFP removes lamin B1 from the region of the nuclear matrix where XY body attaches, possibly through its potential E3 ubiquitin ligase activity. All these hypotheses should be tested in future studies to better understand the underlying mechanisms.

Neither O-glycosylation nor sumoylation of RFP seem to be associated with its specific localization in pachytene spermatocytes. Indeed, RFP (58 kDa) and O-glycosylated RFP (68 kDa) are evidenced at equivalent ratios in meiotic and postmeiotic germ cells (Figure 1(d)). Matsuura et al. [7] have reported that overexpression of SUMO-I in cell culture leads, in somatic cells, to the appearance of a 90 kDa sumoylated RFP. They also have proposed that RFP sumoylation could be involved in the localization of RFP in pachytene spermatocytes. However, we did not detect the described 90 kDa SUMO-I-RFP reactive band on western blot analysis performed on extracts enriched in pachytene spermatocytes (Figure 1(d)). Moreover, our results indicate that SUMO-I and RFP were closely located in spermatocytes but did not colocalize in the XY body (Figures 2 and 3).

The results reported here showed that RFP is able to act as an E3 ubiquitin ligase with a series of E2s such as UbcH1, UbcH5A, 5B, 5C, UbcH6, and HR6B. This activity is exerted by the RING-B-Box domain and requires both ATP and Zn2+. HR6B has been described to be required for correct spermatogenesis and to be localized in the XY body [34, 35]. We propose that a functional interaction between RFP and HR6B could account for an essential role of RFP during spermatogenesis. These hypotheses have to be further investigated in vivo and yeast double-hybrid experiments are currently under way in our laboratory to test a possible interaction between RFP and HR6B.

Like RFP, other E3 proteins such as RNF4, BRCA1, and ICP0 have also been initially described for their broad effect on the cellular transcription activity [28, 32, 33]. It is then interesting to note that, in somatic cells, these proteins, as well as RFP and polyubiquitinated products, colocalize with PML bodies and are able to self-assemble in such structures [12, 25, 36, 37]. We have demonstrated that RFP proteins...
Figure 5: RFP has an E3 ubiquitin ligase activity. (a)–(c) Streptavidin peroxidase revelation of substrate-independent in vitro ubiquitination reactions using immunoprecipitated RFP (a) or GST-RING-B-box of RFP (GST-RING/RFP) (b), (c) as potential E3 ubiquitin ligase enzyme. Dark smears and signals higher than the biggest protein in the assay (monoubiquitinated E1 enzyme indicated by the asterisk) reflected polyubiquitinated products. E2 enzymes are indicated on the top part of each figure. M: master mix containing all the components of the system except E2 enzymes. Sizes (kDa) are indicated. (a) RFP protein immunoprecipitated from 15P-1 cell overexpressing Myc-RFP. Polyubiquitination is observed when UbcH1, UbcH5a, 5b, 5c, UbcH6, or HR6B are used as E2 enzymes. (b) Reactions are performed in presence or in absence of ATP. As expected mono- and poly-ubiquitinations are only observed in presence of ATP. (c) Prior to the substrate-independent in vitro ubiquitination reaction, GST-RING/RFP was pretreated with TPEN (left panel) in order to chelate Zn²⁺ and destroy RING finger architecture, or pretreated with TPEN, washed, and incubated with an excess of Zn²⁺ (right panel). As expected, when GST-RING/RFP was treated with TPEN, only mono-ubiquitination can be observed (left panel) and incubation of TPEN treated GST-RING/RFP with an excess of Zn²⁺ restores E3 ubiquitin ligase activity (right panel).

assemble as a cap in the nuclear envelope matrix, facing the XY body. Later in spermatogenesis, RFP condenses as a spot in round spermatids and these foci are subsequently found in residual bodies, evocating a degradation process (Figure 1). One can postulate that these foci structures result from RFP self-assembly. These observations imply that RFP persists through spermatogenesis and is probably polyubiquitinated and degraded. We propose that RFP acts as an E3 ubiquitin ligase, in addition to its previously described role as transcriptional regulator in somatic cells [17, 18]. One attractive hypothesis could be that RFP, via its E3 ubiquitin ligase activity, might be involved in the release of XY body from the nuclear matrix when pachytene spermatocytes progress through meiotic prophase I. Whether the RFP containing foci structures are the cause or the consequence of XY body release remains to be determined.

It is also interesting to consider the link between PML bodies and XY bodies. We know that both structures share common proteins, for example, PML, DAXX, BRCA1, SUMO-1, and γH2AX, involved in polyubiquitination processes and are sites where transcriptional repression can be established and maintained [25, 31, 38]. So, some of these proteins are able to establish and maintain transcriptional repression by sequestrating transcriptional regulators far away from their targets [25, 38]. Therefore, one might postulate that, in meiotic cells, RFP could interact with transcriptional regulators and that these interactions could play a role in the transcriptional silencing of the XY body through RFP E3 activity.

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