The Potential Mechanism of ZFX Involvement in Cell Growth

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Abstract

Background: The zinc-finger X linked (ZFX) gene encodes a transcription factor that acts as a regulator of self-renewal of stem cells. Due to the role of ZFX in cell growth, understanding ZFX protein-protein interactions helps to clarify its proper biological functions in signaling pathways. The aim of this study is to define ZFX protein-protein interactions and the role of ZFX in cell growth.

Materials and Methods: The PIPs output includes three interacting proteins with ZFX: eukaryotic translation initiation factor 3 subunit I (EIF3I), eukaryotic translation initiation factor 3 subunit G (EIF3G) and protein nuclear pore and COPII coat complex component homolog isoform 3 (SEC13L1).

Results: As a cargo and transmembrane protein interacting with Sec13, eIF3I and eIF3G, ZFX mediates cargo sorting in COPII vesicles at ER exit sites. While traveling to cis-Golgi, eIF3I is phosphorylated by the mechanistic target of rapamycin (mTOR). Proteins transport by COPI vesicles to the nucleouster site layer containing SEC13 via the contribution of microtubules. EIF3G and eIF3I interact with coatomer protein complex subunit beta 2 (COPB2) that helps to enclose ZFX in COPII vesicle. ZFX and eIF3G enter nucleolus where activation of transcription from pre rDNA genes occurs.

Conclusion: We proposed a model in which ZFX is involved in cell growth by promoting the transcription of rDNA genes.

Keywords: ZFX; Cell growth; Cancer stem cells; rDNA genes; Protein-protein interactions

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Introduction

Located in Xp21.3–Xp22.1 and identified as a result of its strong homology to ZFY, the ZFX gene may function in sex determination and escapes X inactivation (1, 2). Differential splicing of ZFX transcript results in five variants. Variants 1, 2 and 3 encode for protein isoform I, variant 4 encodes for isoform II and variant 5 encodes for isoform III. All isoforms encode proteins composed of a highly acidic amino-terminal domain, a basic putative nuclear-localization signal and only isoforms I and II contain carboxy-terminal zinc-finger domain(3). Due to the presence of an acidic domain in ZFX protein, similar to the activation domain of transcription factors, and a potential nucleic acid-binding domain of 13 "zinc fingers", Palmer et al suggested that ZFX may be a plausible transcription factor (4). L’Haridon et al. provided the first demonstration that the ZFX gene encodes a bona fide transcription factor (5). Zinn et al mapped one or more Turner syndrome traits, including short stature, ovarian failure, high-arched palate, and autoimmune thyroid disease in a critical region in Xp11.2-p22.1 (6), implicating the importance of ZFX during game to genes is development and in efficient gametogenesis during
adulthood (7). Of note, Galan-Caridad et al. recently showed that Zfx is a shared transcriptional regulator of embryonic (ESCs) and hematopoietic stem cells (HSCs) (8). Furthermore, Harel et al. reported that ZFX-overexpressing human ESCs produce more colonies with larger size compared to controls (9). Recently, Huang et al. showed that ZFX is up-regulated in cancer stem-like cells in esophageal carcinoma cell lines (10). Furthermore, ZFX is overexpressed in several types of malignancies including diffuse large B-cell and follicular lymphoma (11) and prostate cancer (12). Moreover, it is overexpressed in gastric cancer and our recent work showed a positive correlation between ZFX isoform 3/variant 5 transcript and gastric tumor size, (13, 14) and gliomas. We also found that ZFX gene expression was significantly correlated with central features of the neoplastic phenotype (15, 16). Knocking-down of ZFX in Hep-2, U251 and PC-3 human cancerous cell lines showed that this gene is important in the proliferation and apoptosis of the cells (15). On the other hand, previous studies on ZFX gene has also focused on exploring ZFX target genes and did not analyze its protein-protein interactions. Protein-protein interactions inter-link a huge network of genes and involve in almost every biological process under physiological conditions (17). Due to the role(s) of ZFX in various biological processes such as cell growth, survival, differentiation, cell cycle and apoptosis, understanding ZFX protein-protein interactions helps to clarify its proper biological functions that in turn leads to find its involvement in disease-associated signaling pathways. To this end, the aim of the current study is to define ZFX protein-protein interactions and the role of ZFX in cell growth. We searched PIPs (human protein-protein interaction prediction database) to find ZFX interacting proteins (18).

Materials and Methods

We performed literature mining using various keywords, including “ZFX”, “cancer”, “stem cell”, “transcription factor”. To explore the possible ZFX protein-protein interactions, we referred to several databases including human protein-protein interaction prediction (PIPs) (18), the human protein interaction (HPID) (19), the database of interacting proteins (DIP) (20), and the molecular interaction (MINT) (21). Furthermore, we deduced the functional importance of the predicted ZFX protein-protein interactions using several servers and tools including prediction of cellular role, enzyme class and gene ontology category (ProtFun) (22), PredictProtein (23) ProtParam (24).

Results

Predicted protein-protein interactions

Our data mining in relevant databases to find the ZFX protein-protein interactions did not retrieve any relevant result except for PIPs. As is evident in table 1, the PIPs output includes three proteins: EIF3I (eIF3S2), EIF3G (eIF3S4) and SEC13L1.

Table 1. ZFX interacting proteins as predicted by PIPs. The intensity of the color corresponds to the likelihood of the interaction. The Score column is the predicted score of interaction between the two proteins.

| Protein1 | Protein2 | Name of Protein 2 | Module Scores |
|----------|----------|-------------------|---------------|
|          | ZFX     |                   | Expression    | Orthology | Combined | Transitive | Scores |
| EIF3S2   |          | TRIP1: Eukaryotic translation initiation factor 3 subunit 2 |               |           |          |            | 7.39    |
| EIF3S4   |          | Eukaryotic translation initiation factor 3 subunit 4 |               |           |          |            | 7.39    |
| SEC13L1  |          | N/A               |               |           |          |            | 3.34    |

Discussion

EIF3I and EIF3G are two subunits of eIF3 that binds to the 40S ribosome in an early step of translation initiation and promotes the binding of methionyl-tRNAi and mRNA (25). SEC13 is a Trp- Asp (WD) -repeat protein, localized to the endoplasmic reticulum (ER) exit sites and required for COPII vesicle biogenesis and a shuttle protein between the nucleus and the cytoplasm (26).

Plausible consequences of the protein-protein interactions on post-translational modification (PTM) of ZFX

ZFX, EIF3I and EIF3G genes are transcribed by Ying Yang 1 (YY1) transcription factor (27) and then are translocated to the cytoplasm for translation. The ZFX mRNA may be translated by ribosomes bound to the rough endoplasmic reticulum (RER) since it has an ASN (Asparagine) glycosylation site (28). The
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nascent ZFX protein then may enter the lumen of RER by a doughnut shaped pore consisting of Sec61 protein translocator (29). We postulated that ZFX is then transported to the Golgi apparatus by COPII vesicles enclosed by SEC13-SEC31 heterotetramer for post-translational modification (PTM) such as phosphorylation, the addition of ZN ion and acetylation (28, 30). All of the proteins that need to leave the ER, do not contain a signal for direct binding to Sec24.

After disjoining of the ER exit site to Golgi network, the COPII coat is disassembled by GTP hydrolysis and Sec13-Sec31 cage is depolymerized while SEC23-24 complex is retained (30). The remaining vesicle travels to cis-Golgi for other modifications such as: addition of Zn++, acetylation and amidation (28). While traveling to cis-Golgi, eIF3I is phosphorylated by mammalian target of rapamycin (mTOR) (36) localized to the endoplasmic reticulum and Golgi (35). Transported proteins from Golgi to the nucleus are carried by COPI vesicles to nuclear pore complex containing SEC13 in outer site layer (37) via the contribution of microtubules (38). The coat structure of COPI vesicles consists of the heptameric coat protein complex that COPB2 (beta'-COP) (39) protein is a part of this complex (40). As predicted by PIPs, eIF3G and eIF3I interact with COPB2 (Figure 2).

Therefore, we postulated that this interaction may help to enclose ZFX in COPI vesicles (41) (Figure 3). As eIF3I contains the WD-40 domain, it is plausible that it functions as an adaptor protein (42). COPI vesicle coat proteins are disassembled by hydrolysis of GTP (38) and lyase enzymatic property of eIF3I (22) may result in the breakage of (N-C) link that binds myristic acid in the vesicle membrane to a nitrogen of an amino acid in eIF3I protein, therefore eIF3I protein disassociates from eIF3G and ZFX proteins. eIF3I is a scaffold protein for assembly of eIF3 factor subunits (42) in which bound eIF3

Figure 1. Cargo (ZFX) sorting mediating by the adaptor protein (SEC13) in COPII vesicles at ER exit sites (modified from Zanetti et al. 2011). The ZFX mRNA may be translated by ribosomes bound to the rough endoplasmic reticulum (RER); the nascent ZFX protein then may enter the lumen.

Some of these proteins might interact with a transport adaptor like SEC13 (containing a WD repeat region) (22), and thus be included in the COPII vesicle through an indirect interaction (31). Due to the presence of an NLS (nuclear localization signal) domain in ZFX protein (32), it can be regarded as a cargo. This cargo may interact with SEC13 (33) as predicted by PIPs (Table 1). Furthermore, the presence of an N-myristoylation site and a beta barrel structure in ZFX protein implicates that it is a transmembrane protein (TMP) (22). Being as a cargo and transmembrane protein interacting with Sec13, ZFX might interact with eIF3I and eIF3G (as cytoplasmic accessory proteins which are adaptor proteins involving in signal transduction) to mediate cargo sorting in COPII vesicles at ER exit sites. As the size and function of a COPII vesicle coat are regulated by ubiquitin (34), and ZFX, eIF3I, eIF3G and SEC13 proteins interact with ubiquitin C (UBC) (35), we postulated that the above-mentioned process of formation of ZFX-containing COPII vesicles may be plausible (Figure 1).
inhbts binding of 60S subunit to 40S subunit and promotes the recruitment of the 40S ribosomal subunit to the mRNAs (25) and therefore remaining of the Elf3l near NPC is rational. However, Elf3g possesses two RNA binding domains (RBS) (43); therefore, it may function in transcription regulation as predicted by ProtFun 2.2 server (22). Moreover, it can bind to 18S rRNA (44). Taken together, it is possible that ZFX and Elf3g enter the nucleolus where they activate transcription from pre rDNA genes. The involvement of a zinc finger protein in rRNA gene transcription has been reported for Basonuclin, a zinc finger protein of keratinocytes (45). Furthermore, there are multiple zinc finger recognition sites in the promoter of rDNA genes as predicted by zinc finger tools (46). Besides, due to the possible dual role of Elf3g in transcription regulation and binding to the transcript of rDNA genes to prevent them from the assembly of ribosome; we proposed a model in which ZFX and its interacting proteins including Elf3g contribute to cell growth by promoting the transcription of rDNA genes in the nucleolus.

**Figure 3.** Schematic representation of the assembly of COPI vesicle containing ZFX (modified from Bethune et al. 2006).

**Conclusion**

In our model, we proposed that ZFX may interact with Elf3l, Elf3g and Sec13l1. After transcription, the ZFX mRNA may enter the lumen of the rough endoplasmic reticulum by Sec61. Being as a cargo and transmembrane protein interacting with Sec13, ZFX might interact with Elf3l and Elf3g to mediate cargo sorting in COPII vesicles to transfer to the Golgi apparatus for PTM. Interaction between Elf3g, Elf3l and COPB2 may help to enclose ZFX in the COPII vesicles in the Golgi apparatus. Once COPII vesicles disassemble, the Elf3l may remain near the NPC, while ZFX and Elf3g may enter the nucleus where they activate transcription from pre rDNA genes.

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**Author Contributions**

All authors made substantial contributions to conception and design, acquisition of data, analysis and interpretation of data; all authors participated in drafting the article or revising it critically for important intellectual content. All authors gave final approval of the version to be submitted in any revised version.

**Conflict of Interest**

Authors declare no conflict of interest.

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