Unexpected distribution of KRIT1 inside the nucleus: new insight in a complex molecular pathway

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Abstract

KRIT1 is an 84kDa protein that lacks any relevant catalytic domains, associated with the cerebral cavernous malformation disease. We have investigated by means of ultrastructural immunocytochemistry the nuclear distribution of KRIT1 in different cell lines, revealing its unexpected localization on actively transcribing nuclear domains such as the perichromatin fibrils and the nucleolar dense fibrillar component. These preliminary data indicate a still undescribed and unknown role for KRIT1 inside the nucleus.

Introduction

KRIT1 (Krev1 interaction trapped gene 1) is a gene associated with the cerebral cavernous malformation (CCM) disease, characterized by vascular abnormalities that cause risk of hemorrhages in the brain with consequent headache, seizures or even death. Along with KRIT1, two other genes, CCM2 (cerebral cavernous malformation 2) and PDCD10 (programmed cell death 10), were linked to CCM disease, all the three genes coding for scaffolding proteins lacking catalytic domains but presenting several domains for protein-protein interaction. In the last years, several studies have provided new information on the signaling pathway in which KRIT1 takes part, suggesting a role in vascular development, cell-cell junctions, and cell-extracellular matrix adhesion. However, none of these studies was focused on the nuclear compartment where it is known KRIT1 can be physiologically present, as seen by fluorescence resonance energy transfer (FRET) experiments and immunofluorescence. Indeed, KRIT1, together with its protein interacting partner, ICAP1 (integrin beta-1-binding protein 1), shuttle between the nucleus and the cytoplasm bearing a putative NLS in its N-terminal amino acid sequence.

Although the shuttling dynamics involving KRIT1, ICAP1 and CCM2 is only partially understood, there is no evidence in literature of a possible active function of KRIT1 inside the nuclear compartment. One of the current hypotheses is that KRIT1 and ICAP1 regulate each other by sequestering the partner inside the nucleus, thus preventing interaction with cytoplasmic proteins, such as beta-1 integrin or the p-38-MAPK associated complex. By means of ultrastructural immunocytochemistry we detected KRIT1 inside the nucleus with the aim to gain insight into other possible roles of this protein, as suggested by its localization on nuclear structures of known function. The nuclear location of KRIT1 was investigated by transmission electron microscope in HeLa cells under normal growth conditions or after heat shock when the nuclear distribution of ribonucleoproteins (RNP)s is modified, as well as in rat hepatocytes and in Chironomus thummi salivary glands (the latter being an excellent model for transcription due to the presence of actively transcribed polytenic chromosomes).

The results obtained suggest that KRIT1 is involved in the maturation of nascent RNA transcripts, due to its prevalent localization in nuclear sites where active transcription takes place.

Materials and Methods

Cells and treatments

HeLa cells were grown in DMEM medium supplemented with 10% fetal bovine serum, 2mM glutamine and 100 units each of streptomycin and penicillin, in 25 cm2 plastic flasks or multi-well dishes (recognizing the 3mG cap of snRNAs, kindly provided by Dr. R. Lührmann) diluted 1:5; anti H20 tapeptide epitope of RNA Polymerase II, and anti P70 in PBS containing the primary anti-KRIT1 antibody (rabbit-polyclonal against human epitope of KRIT1, GenTex, GTX109560) diluted 1:5, and eventually one of the following primary mouse monoclonal antibody to detect colocalization: anti-Pol II (against human hep-tapeptide epitope of RNA Polymerase II, and recognizing both the active and inactive forms; Biodiesign, M55701M) diluted 1:5; anti H20 recognizing the 3mG cap of snRNAs, kindly provided by Dr. R. Lührmann) diluted 1:5. After rinsing with PBS-Tween and PBS, grids were incubated with NGS as above, and then reacted with a goat-anti rabbit and a goat anti-mouse secondary antibodies conjugated with 12 nm or 6 nm colloidal gold particles (The Jackson Lab., Bar Harbor, ME, USA) for 30 min at room temperature, rinsed and finally stained. To reduce chromatin contrast and selectively reveal nuclear ribonucleoprotein constituents, the sections were bleached by the EDTA method or selectively stained for RNA with terbium citrate.

The samples were observed on a Zeiss 900 EM operating at 80kV. In some of the micrographs, the gold grains have been digitally contrasted with the Photoshop software.

Results

After EDTA staining to reveal RNPs, KRIT1 labeling in HeLa cells was localized on the perichromatin region, where transcription takes place and, in particular, on the perichromatin fibrils (PF, the in situ morphological expression of the extranucleolar RNA tran-
This pattern was also observed after heat shock, suggesting that the nuclear localization of KRIT1 is not influenced by this type of stress. More in detail, staining with terbium citrate for RNA allowed us to better visualize the nascent PFs labeled with the KRIT1 antibody (Figure 1b).

We were able to observe the same pattern in salivary glands of *Chironomus thummi*: Fig 1c shows a PF where the labelings for RNA Pol II and KRIT1 are present at the opposite extremities. Since the antibody does not discriminate between the active and inactive form of RNA Pol II, we cannot determine if that PF is being transcribed, although it is tempting to assume so. In this model system, KRIT1 is also present in the large (perichromatin-granules like) Balbiani granules, where the mRNA transcribed is stored for the export to the cytoplasm (Figure 2a). We rarely found KRIT1 co-localizing with RNA Pol II, whereas we observed a co-localization with the methyl-cap of snRNA. In Figure 2b an example of double labeling in a HeLa cell after heat shock is presented. In all the cell types used in these experiments, we have observed focal electron dense areas of the nucleus that show a higher labeling for KRIT1 suggesting the presence of nuclear bodies of unknown function (Figure 2 c,d). These labeled nuclear domains were observed in all immunolabeling experiments in condition of minimum background, which leads us to exclude an artifactual labeling. Labeling for Pol II was never found on these nuclear domains.

Finally, we have documented KRIT1 presence in the dense fibrillar component (DFC) of the nucleolus in numerous HeLa samples (Figure 2 e,f; Supplementary Figure 2). The same labeling distribution is present on the nucleolus in hepatocytes (not shown).

**Discussion**

Our results provide novel information about the fine nuclear distribution of the protein KRIT1 and its spatial relationships with the RNP components involved in RNA transcription and maturation, confirming and extending previous data on its nuclear localization.

Despite its known role in cell-cell junction and cell-matrix anchorage, KRIT1 shows a peculiar nuclear labeling on the PFs and nucleolar DFC, where transcription actively takes place. The recurrent presence of this labeling pattern in most of the cells in the samples observed, from human HeLa cells to

**Figure 1.** a) KRIT1 labeling pattern at low magnification; EDTA staining; the gold particles (highlighted in red) are present on the PF at the periphery of condensed chromatin areas (ch) and on the nucleolus (Nc); scale bar: 500 nm. b) A nascent single PF stained with terbium for RNA in a HeLa cell nucleus; the large gold grain represents RNA Pol II while the small grain indicates KRIT1; scale bar: 100 nm. c) The same labeling pattern as in (b) was observed in a *Chironomus* nascent fibril; scale bar: 100 nm.

**Figure 2.** a) Four Balbiani granules are shown, one of which is labeled for KRIT1 at the periphery; terbium staining; scale bar: 50 nm. b) HeLa cell double labeled for m3Gcap (small grains) and KRIT1 (large grains); in some cases (arrow) we can observe a co-localization; EDTA staining; scale bar: 100 nm. c) Rat liver after EDTA staining; a nuclear domain is shown with a strong labeling for KRIT1; scale bar: 100 nm. d) A similar nuclear domain, terbium stained, is present in a HeLa cell; also in this case a heavy labeling for KRIT1 is present; scale bar: 100 nm. e) Labeling for KRIT1 in the nucleolus of a HeLa cells; the grains are present on the dense fibrillar component; scale bar: 200 nm. f) Part of a nucleolus of a Hela cell nucleus; KRIT1 is present in the dense fibrillar component surrounding the fibrillar center (Fc); terbium staining; scale bar: 50 nm.
Chironomus thummi salivary glands, suggests a possible conserved function of KRIT1 inside the nuclear compartment, reasonably involving RNA transcription. The presence of the labeling at the terminus of the newly transcribed PFs (i.e., far away from Pol II but associated with the m3G cap), together with the lack of DNA binding domains in the KRIT1 protein structure, leads us to hypothesize that KRIT1 could be involved in RNA maturation rather than be directly involved in transcription. Also the presence of KRIT1 labeling in the perichromatin-granules-like Balbiani granules could support this idea.

No marked difference was observed in Hela cells before and after heat shock, suggesting that KRIT1 nuclear localization and/or shuttling is not modified or increased by heat shock. Only few cells within the various samples studied showed the high labeled nuclear domains. As it is the case for other nuclear bodies, their function could be related to the nucleus-cytoplasmic shuttling of KRIT1. Being positive to Terbium and EDTA staining we know they are composed of RNPs, but further studies are needed to understand the nature of these nuclear domains.

These data lay the basis for a further characterization of KRIT1 function inside the nuclear compartment. It is not excluded that an impairment of this unknown KRIT1 nuclear function could contribute to the development of CCM in the individuals that carry mutations on this gene. Thus we expect an altered KRIT1 labeling pattern in CCM patients relative to healthy. Furthermore, these data can be implemented by observing the nuclear labeling pattern of the CCM2 and PDCD10 protein products, involved in the development of the disease. Being the pathogenetic mechanisms by which KRIT1 mutation cause familial CCM still unknown, we believe that adding new information on the complex molecular pathway of these genes could lead to a better understanding of the CCM pathology.

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