# Rev A/2013/03/12/SF-797DSCA/AK

# TIMP2 Hu-Cy5 SmartFlare<sup>TM</sup> RNA Detection Probe

Cat. # SF-797

FOR RESEARCH USE ONLY

pack size: 50µL (250 rxns)

Store at 2-8°C, after reconstitution store at 23-27°C DO NOT FREEZE



## **Product Data Sheet**

page 1 of 1

Validated Accession NM 003255.4

NOT FOR USE IN DIAGNOSTIC PROCEDURES

NOT FOR HUMAN OR ANIMAL CONSUMPTION

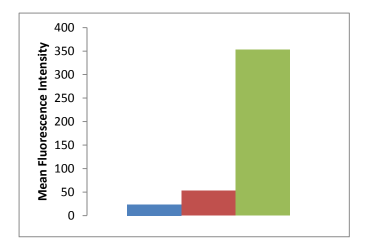
Species Hu

Gene Aliases CSC-21K

### Confirmation of TIMP2 SmartFlare Performance:

TIMP2 SmartFlare has been tested in a buffer system to detect the release of the fluorophore in the presence of a complementary base pair sequence for each lot to confirm target specificity.

TIMP2 SmartFlare has also been tested in a cell model system and demonstrated increased fluorescence in cells expressing the target compared to a scrambled negative control SmartFlare (Figure 1). For additional accession numbers predicted to react with this SmartFlare Probe, please visit http://www.millipore.com/catalogue/item/SF-797



**Figure 1:** TIMP2 Mean Fluorescence Intensity (green) measured by flow cytometry in living MDA-MB-231 cells demonstrated a significant increase over unflared cells (blue) as well as scramble control (red). Data shown in graph is representative.

### Storage and Handling:

Material has been 0.22µm filtered. Stable for 5 years at 2-8℃ degrees in lyophilized format ONLY. Room temperature is required for reconstituted product.

Warning-after reconstitution product is sensitive to cold and hot temperatures, a stable room temperature of 23-27°C is required.

### **Handling Recommendations:**

Reconstitute with sterile nuclease free water in a drop wise fashion and tap tube repeatedly to fully dissolve lyophilized material. Vortex for 5-10 sec.

Upon reconstitution, store at room temperature for up to 1 year protected from light. Product must be handled with gloves as product can be absorbed through the skin.

### **Recommended Cell Testing Protocol:**

(example: 30,000 cells in a 200µL media volume within each well of a 96 well plate)

- Reconstitute reagent in 50µL of sterile nuclease free water.
- Create a working solution based on your experiment by diluting 1:20 in sterile PBS.
- Add 4µL directly to cells (at approx 80% confluency).
- Allow to incubate overnight for 16 hrs.
- Detect using fluorescence detection platform of choice.

