A GFP-based reporter system to monitor nonsense-mediated mRNA decay

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ABSTRACT

Aberrant mRNAs whose open reading frame (ORF) is truncated by the presence of a premature translation-termination codon (PTC) are recognized and degraded in eukaryotic cells by a process called nonsense-mediated mRNA decay (NMD). Here, we report the development of a reporter system that allows monitoring of NMD in mammalian cells by measuring the fluorescence of green fluorescent protein (GFP). The NMD reporter gene consists of a T-cell receptor-β minigene construct, in which the GFP-ORF was inserted such that the stop codon of GFP is recognized as PTC. The reporter mRNA is therefore subjected to NMD, resulting in a low steady-state mRNA level, an accordingly low protein level and hence a very low green fluorescence in normal, NMD-competent cells that express this reporter gene. We show that the inactivation of NMD by RNAi-mediated knockdown of the essential NMD factor hUpf1 or hSmg6 increases the NMD reporter mRNA level, resulting in a proportional increase of the green fluorescence that can be detected by flow cytometry, spectrofluorometry and fluorescence microscopy. With these properties, our GFP-based NMD reporter system could be used for large-scale screenings to identify NMD-inhibiting drugs or NMD-deficient mutant cells.

INTRODUCTION

Quality control checkpoints at different stages during gene expression are necessary to keep production of aberrant gene products low and thus to allow proper functioning of the cells. On the post-transcriptional level, a process called nonsense-mediated mRNA decay (NMD) has evolved in all eukaryotes examined so far to recognize and specifically degrade aberrant mRNAs, in which the open reading frame (ORF) is truncated by the presence of a premature translation-termination codon (PTC) (1). Two important sources giving rise to PTC-containing mRNAs (PTC+ mRNAs) in metazoans are mutations in the DNA and pre-mRNA splicing. It is estimated that about one-third of all inherited genetic disorders and many forms of cancer are caused by mutations that result in the generation of PTC+ mRNA (2). In humans, more than half of all pre-mRNAs undergo alternative splicing (3), and about one-third of these alternative transcripts contain a PTC (4). This demonstrates that NMD is an important cellular process that prevents (or at least reduces) the production of potentially deleterious truncated proteins and thereby modulates the clinical manifestations of many genetic diseases (2,5,6). In addition, immunoglobulin and T-cell receptor (TCR) genes more often than not acquire PTCs during the programmed V(D)J rearrangements which take place during lymphocyte maturation (7).

The molecular mechanisms underlying NMD are not well understood. Seven different NMD factors have been genetically identified in Caenorhabditis elegans (8,9), and the orthologues of three of them have also been genetically identified in Saccharomyces cerevisiae (10,11). The orthologue genes in humans, mice and Drosophila have all been found by homology searches (12–20). Owing to the lack of functional screens, putative mammalian NMD factors without orthologues in a genetically amenable organism are therefore still awaiting their discovery. It seems likely that additional, yet unknown, NMD factors might exist in vertebrates, considering that NMD appears to be more regulated and more sophisticated in more complex organisms. For example, hUpf1 activity appears to be regulated by a cycle of phosphorylation and dephosphorylation in metazoans, while S.cerevisiae lacks the orthologues involved in regulating hUpf1’s phosphorylation state (1). Of interest for this study, the human orthologue of C.elegans SMG6, hSmg6, functions in dephosphorylation of hUpf1 (20) and is required for NMD (19). In addition, hSmg6 was also found to associate with telomerase and its overexpression

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uncaps telomeres (18). Also consistent with the idea of more factors being involved in NMD of more complex organisms, several factors of the exon junction complex were found to play a role in human NMD (21–25), while in Drosophila, the exon junction complex is dispensable for NMD (19).

Although technically quite demanding, mutagenesis screens in diploid human cells have been shown to be feasible (26,27), provided a suitable reporter system is available. To study mammalian NMD, several genes with PTCs at various positions were established as ‘model systems’ (28–32). But since the analysis of NMD of these genes requires analysis of mRNA, they are not suitable as reporters for screens. We, therefore, decided to develop a NMD reporter system where the readout was based on fluorescence in intact cells or whole cell extracts. We figured that such a NMD reporter system would not only be a valuable tool for basic research but could also be useful for drug screening. Given the large number of genetic diseases originating from PTC-causing mutations, there is also a considerable pharmaceutical interest to develop substances that interfere with NMD. Here, we describe the construction and characterization of a green fluorescent protein (GFP)-based NMD reporter system that allows detection of NMD-deficient cells by the detection of increased green fluorescence. We show that this NMD reporter system can be used in conjunction with flow cytometry, spectrofluorometry and fluorescence microscopy.

**MATERIAL AND METHODS**

**Plasmids**

The parental vector expressing the TCR-β minigene (pβ433) has been described in detail elsewhere (33). pβ510 was generated from pβ433 by replacing the NotI–SalI fragment with a double-stranded oligo (5’-ggGCGCGCCGCGCGCGCTTATAAAAA-CCACGGGAGACGGGACGCAACCACCCGCCAGACCCGACGTACCAC-3’; sequence of forward oligo is in upper case, NotI and SalI sites in italics, transcription start site underlined), which creates an intronless transcribed region of 62 bp. The EGFP ORF was PCR amplified using primers 5’-ATCTATAGCATATGCGATGTGAGACAAGGCGAG-3’ and 5’-ATCGACACCATAATGCTCAGTTACGATCAGCTACAGCAGC-GTCCCATGC-3’ (Ndel site in italics, stop codon underlined), digested with Ndel and inserted into the Ndel site of pβ510.

Finally, the leader exon sequence was exchanged for a sequence encoding the haemagglutinin A epitope by replacing the SalI–EcoRI fragment with a double-stranded oligo (5’-TCGACACCATGCGGCTACCATACTATGATTCCCAGAT-TTACGTCTCACCATGCGAAGCTGATGCTGCTGAGCCACTGTGTTGTCTTTTTTTTGaattc-3’; sequence of forward oligo is in upper case, SalI and EcoRI sites in italics, translation start site underlined). The resulting plasmid is called pβ510-HA-TCRβ-GFP. The plasmid pβ510-HA-TCRβ-GFP.JCin was generated by deleting the JC intron using fusion PCR. Detailed cloning strategy and primer sequences are available upon request. The HA-TCRβ-GFP gene of both plasmids was sequenced.

pSUPERpuro vector was created by inserting a puromycin-resistance cassette lacking the HindIII site of the puromycin ORF (from pTRE2pur, Clontech) into the Xhol site of pSUPER (34). Sequences coding for short hairpin RNAs (shRNAs) were inserted as double-stranded oligos into pSUPERpuro between the BglII and HindIII as described previously (34). The two target sequences of hUpf1 were 5’-GAGAATGCCTACTTACACT-3’ (pSUPERpuro-hUpfI) and 5’-GATGCGAGTCCCCTCATT-3’ (pSUPERpuro-hUpfII) and the two target sequences of hSmg6 were 5’-GGTCACAGTGCTGAAGTA-3’ (pSUPERpuro-hSmg6/I) and 5’-GCTGCAAGTTACTTACAAG-3’ (pSUPERpuro-hSmg6/II).

**Cell culture and transfection**

HeLa cells were grown in DMEM (Invitrogen) supplemented with 10% heat inactivated fetal calf serum (FCS), 100 U/ml penicillin and 100 μg/ml streptomycin. LipofectAmine (Invitrogen) was used for all transient transfections according to the manufacturer’s protocol. For RNAi experiments, cells were seeded in 8 cm dishes and transfected with 2 μg empty pSUPERpuro (mock), a mixture of 1 μg pSUPERpuro-hUpfI and 1 μg pSUPERpuro-hUpfII, or a mixture of 1 μg pSUPERpuro-hSmg6/I and 1 μg pSUPERpuro-hSmg6/II using 8 μl LipofectAmine (Invitrogen). An aliquot of 1.5 μg/ml puromycin (Calbiochem) was added to the cells from 24 h post-transfection until the evening before analysis (4 days post-transfection) to eliminate the untransfected cells. Before analysis, the cells were cultured without puromycin for at least 16 h to avoid potential effects of this translation inhibitor on NMD. In cells transfected with the empty pSUPER-puro, relative levels of NMD substrates and mRNAs without a PTC were identical irrespective of whether the cells were treated with puromycin or not (data not shown), confirming that the puromycin treatment did not interfere with NMD. For microscopy, 10 mm glass coverslips were placed in the culture dish before seeding the cells.

To establish NMD reporter-expressing HeLa cell lines, pβ510-HA-TCRβ-GFP PTC+ was transfected with Gene Jammer Transfection Reagent (Stratagene) according to the manufacturer’s instructions, and stable transfectants were selected by cultivating the cells in the presence of 500 μg/ml of G418 (Geneticin, Invitrogen) for 2 weeks, before single cell clones were isolated and expanded. As a control, stably transfected clonal cell lines were also established using a reporter construct that lacks the JC intron (pβ510-HA-TCRβ-GFP.JCin).

The monoclonal cell lines were treated with 100 μg/ml cycloheximide (CHX) for 3 h to analyse the response of NMD reporter mRNA or mRNA of the ΔJCin control construct to inhibition of NMD.

**RNA analysis**

Total cellular RNA was purified using the Absolutely RNA RT–PCR Miniprep Kit (Stratagene) according to the manufacturer’s protocol. For real-time PCR analysis, 1 μg was reverse transcribed in 50 μl Stratascript buffer in the presence of 200 ng random hexamers, 0.4 mM dNTP mix, 40 U RNaseIn (Promega) and 50 U Stratascript (Stratagene) according to the manufacturer’s protocol. Reverse transcribed material corresponding to 40 ng RNA was amplified in 25 μl Universal PCR MasterMix, no AmpErase (Applied Biosystems) using the following primers and TaqMan probes. TCR-β mRNA was measured over the junction between the VDJ exon and the C1 exon using 400 nM forward primer om59 (5’-GCAGCTGCAGGTTACTTACAAG-3’), 400 nM reverse primer om60...
(5’-TGGCTCAAACAAGGAGACCTT-3’) and 200 nM TaqMan probe TM59/60 (5’-FAM-CTCGAGGATCTGAGAAATGTGACTCCACC-TAMRA-3’). β-Globin mRNA was measured over the junction between exons 2 and 3 using 800 nM forward primer om111 (5’-GCTGCACGTGACGAAGCTGC-3’), 800 nM reverse primer om112 (5’-AAAGTGATGGGGCAACGAC-3’) and 200 nM TaqMan probe TM111/112 (5’-FAM-TCCTGAGAACTTCAGGCTCCTGGCAAC-TAMRA-3’). β-Globin mRNA was measured over the junction between exons 2 and 3 using 800 nM forward primer om111 (5’-GCTGCACGTGACGAAGCTGC-3’), 800 nM reverse primer om112 (5’-AAAGTGATGGGGCAACGAC-3’) and 200 nM TaqMan probe TM111/112 (5’-FAM-TCCTGAGAACTTCAGGCTCCTGGCAAC-TAMRA-3’). hUpf1 mRNA, GAPDH mRNA and 18S rRNA were measured using pre-developed assay reagents from Applied Biosystems. For 18S rRNA measurements, cDNA corresponding to 400 pg RNA instead of 40 ng was used. Real-time PCR was run on the GeneAmp 5700 and 7000 Sequence Detection Systems (Applied Biosystems) using the standard thermal profile.

The conditions for the traditional endpoint RT–PCR shown in Figure 1B were as follows: RNA was reverse transcribed using an anchored primer against the poly(A) tail (5’-N30-VN-3’) and Stratascript reverse transcriptase in Stratascript first strand buffer according to the manufacturer’s protocol (Stratagene). Subsequent PCR was performed with a primer binding to the HA-tag (om153; 5’-TCCAGATTACGCTCACTCGAAC-3’) and a reverse primer binding to exon C1 (om96; 5’-GATGGCTCAAACAAGGAGACCTT-3’). An aliquot of the PCR was analysed on a 1.5% agarose gel and stained with ethidium bromide.

For the northern blot shown in Figure 1C, 10 μg of total cellular RNA was separated on a 1.2% agarose gel containing 1% formaldehyde and 1× MOPS. A picture of the ethidium bromide stained gel before blotting was taken to use the 18S rRNA band as loading control. The RNA was transferred to positively charged nylon membrane (Roche Diagnostics) in 20× SSC by standard capillary blotting method. Probe labeling, hybridization, washing and signal detection was as described elsewhere (32). Approximately 50 ng of a purified Sall–BamHI fragment comprising the TCR-β minigene sequence was used as template for the 32P-labelled probe.

**Flow cytometry analysis**

About 10^6 cells were harvested and resuspended in phosphate-buffered saline (PBS)/1% FCS and then analysed using a FACScan (BD, San Jose, CA). GFP fluorescence was excited at 488 nm and emission was measured with a 530/30 nm bandpass filter. HeLa cells without the NMD reporter gene (plain HeLa) served as a control for autofluorescence and were used to set the gate for GFP positive cells. The data were processed using FlowJo software (Tree Star, Ashland, OR).

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**Figure 1.** Expression of the HA-TCRβ-GFP reporter gene yields correctly spliced mRNA. (A) Schematic representation of the NMD reporter gene. Sequences derived from human β-actin, mouse TCR-β and EGFP are shown in white, yellow and green, respectively. The haemagglutinin epitope (HA) is shown in purple. The positions of the transcription start site (+1), translation start codon (ATG), PTC and the normal stop codon (TGA) are depicted, and relevant restriction sites are indicated. The intron deleted in the control construct ΔJCin is marked. See text for details. (B) RNA from HeLa cells transiently (tr) or stably (st) expressing the HA-TCRβ-GFP reporter gene (PTC+ or ΔJCin) and a reverse primer binding to exon C1 (om96; 5’-GATGGCTCAAACAAGGAGACCTT-3’). An aliquot of the PCR was analysed on a 1.5% agarose gel and stained with ethidium bromide.

For the northern blot shown in Figure 1C, 10 μg of total cellular RNA was separated on a 1.2% agarose gel containing 1% formaldehyde and 1× MOPS. A picture of the ethidium bromide stained gel before blotting was taken to use the 18S rRNA band as loading control. The RNA was transferred to positively charged nylon membrane (Roche Diagnostics) in 20× SSC by standard capillary blotting method. Probe labeling, hybridization, washing and signal detection was as described elsewhere (32). Approximately 50 ng of a purified Sall–BamHI fragment comprising the TCR-β minigene sequence was used as template for the 32P-labelled probe.

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The geometric mean values were used to compare the amount of fluorescence in the different samples.

**Western blot analysis**

Cells were lysed and denatured in SDS loading buffer and these whole cell lysates were separated on 10% SDS–PAGE. Proteins were then transferred to Optitran BA-S 85 reinforced nitrocellulose (Schleicher & Schuell) and probed with different antibodies. The monoclonal anti-HA mouse antibody (BabCO, Berkeley, CA) was used at a dilution of 1:1000, the anti-hUfp1 rabbit antiserum ([14], a kind gift from Jens Lykke-Andersen) was diluted 1:2500, and supernatant of the hybridoma cell line Y12 (producing the anti-Sm antibody) was diluted 1:400. Horseradish peroxidase-conjugated anti-mouse or anti-rabbit IgG antibody (Promega) was used as secondary antibodies at a dilution of 1:4000. ECLplus Western Blotting Detection System (Amersham Pharmacia Biotech).

**Dual scanning microplate spectrofluorometer**

Proteins were extracted in 1% deoxycholic acid, 1% NP40, 25 mM Tris–HCl (pH 8), 50 mM NaCl and protease inhibitor cocktail (Complete Mini, EDTA-free; Roche). The cells were incubated at 4°C for 30 min on a shaker and then centrifuged at 16 000 g for 5 min. The supernatant was immediately frozen in liquid nitrogen and stored at −20°C. GFP fluorescence was measured on a SPECTRAMax Gemini (Molecular Devices) using 96-well black plates with flat, black bottom. GFP fluorescence was excited at 488 nm and measured at 512 nm, the cut-off was set at 495 nm. SOFTmaxPRO software was used for data analysis.

**Confocal microscopy**

Images were collected on a Leica TCS SP2 AOBS laser scanning confocal microscope equipped with a HCX PL APO lb.d.BL, 63.0x 1.2W objective (Leica Microsystems Inc., Exton, PA). The cells were grown on coverslips, washed once with PBS and then fixed in 4% paraformaldehyde and mounted on DAKO fluorescent anti-fading mounting medium (DAKO Corporations). For GFP detection, cells were scanned using the 488 nm laser line and emitted light between 500 and 600 nm was collected. The pinhole was opened to 418 μm and the photomultipliers were set to 580 V. The same settings for image acquisition and processing have been applied for all samples to allow comparison of the fluorescence intensities among the different samples.

**RESULTS AND DISCUSSION**

**Design of the NMD reporter gene**

Fluorescent proteins have become a valuable tool for innumerable applications in modern biology, because the light emitted by these proteins can be detected relatively easily by many different techniques. Therefore, we decided to develop a reporter system for NMD that is based on the enhanced green fluorescent protein (EGFP, Clontech). To this end, we inserted the EGFP ORF in frame into the middle exon of a TCR-β minigene (Figure 1A). In this position, more than 50 nt upstream of the 3′-most 5′-splice site, the stop codon of the EGFP ORF is recognized as a PTC and, thus, the mRNA is subjected to NMD. A TCR-β minigene was chosen for our reporter gene, (i) because NMD of TCR-β transcripts has been extensively studied and (ii) because PTCs in these TCR-β minigenes cause a dramatic reduction in the steady-state mRNA levels (35,36). The particular TCR-β minigene used here has fused the first 120 bp of exon C1 to base pair number 117 of exon C4 and, therefore, lacks part of the constant region. Since the signal peptide encoded by the Vleader exon is still present in this TCR-β minigene, while the region of the gene encoding the membrane-spanning domain is missing, it was likely that this TCRβ-GFP protein would be secreted from the cells. To prevent this potential problem, and to introduce a tag that allows easy detection of the TCRβ-GFP protein in western blots, we replaced the signal peptide encoding sequence in the Vleader exon with a sequence encoding for the haemagglutinin epitope (Figure 1A, HA-tag).

In addition, a control construct was derived from the NMD reporter gene by deleting the second intron (Figure 1A, ΔJCin). This ΔJCin construct encodes the same TCRβ-GFP fusion protein as the NMD reporter construct, but because the PTC in the ΔJCin construct is now located in the last exon, its mRNA is not expected to be a substrate for NMD.

In RNA samples from HeLa cells stably or transiently expressing the TCRβ-GFP reporter gene, we amplified by RT–PCR a product of the size expected for the correctly spliced mRNA (Figure 1B), indicating that insertion of the EGFP ORF into the VDJ exon, which results in an unusually long internal exon, did not induce alternative splicing of the pre-mRNA. Correct splicing was also confirmed for the ΔJCin construct stably expressed in HeLa cells (Figure 1B), and for the NMD reporter gene stably expressed in the diploid human cell line HT1080: northern blotting revealed a single and specific band for HA-TCRβ-GFP mRNA in the expected size range (Figure 1C). We conclude that the vast majority of HA-TCRβ-GFP transcripts are spliced correctly in human cells, both when expressed episomally (transient transfection) or from a genomic locus (stable transfection).

The **GFP fluorescence is proportional to the mRNA level of the reporter gene**

The concept of a GFP-based NMD reporter system is to use the GFP fluorescence as a measure for the steady-state mRNA level of the NMD reporter. Thus, the reporter system is only useful, if the intensity of the GFP fluorescence is proportional to the mRNA level. To test this, HeLa cells were transfected with increasing amounts of NMD reporter plasmid DNA. Seventy-two hours post-transfection, a fraction of the cells was analysed by flow cytometry (Figure 2A), and from the rest of the cells, RNA was isolated and relative reporter mRNA levels were measured by real-time RT–PCR. The comparison between the percentage of GFP-positive cells multiplied by the average intensity of GFP fluorescence (Figure 2B) and the relative reporter mRNA levels (Figure 2C) for the different plasmid DNA amounts shows that the GFP signal is indeed highly proportional to mRNA levels.

The same experiment was also carried out with the ΔJCin control construct, and the respective GFP signals (mean fluorescence intensity multiplied by percentage of GFP-positive cells) are shown in comparison with the GFP signals determined for the PTC+ construct (Figure 2D). With increasing
amounts of transfected plasmid DNA, the GFP signal of the NMD reporter construct changes from 9-fold lower than the GFP signal of the NMD-resistant control construct to only 3-fold lower. The extent of NMD cannot be determined accurately from this experiment for two reasons. When low amounts of plasmid DNA is transfected, most of the resulting weak GFP signals will be hidden within the autofluorescence of the cells, and only a small percentage of the cells qualify as 'GFP signals will be hidden within the autofluorescence of the cells, and only a small percentage of the cells qualifies as 'GFP positive' (fluorescence signal distinct from the autofluorescence seen in the mock-transfected control cells) and contributes to the overall GFP signal. Because this effect is more pronounced for the PTC+ than for the ΔJCin, the extent of NMD will be overestimated with low plasmid DNA amounts. On the other hand, we know from experience with a number of different NMD reporter genes that NMD can easily be saturated by the expression of high amounts of NMD substrate (A. Paullusson and O. Mühlemann, unpublished data). This partially saturated NMD results in a decreased difference between the PTC+ and the ΔJCin GFP signal measured with high plasmid DNA amounts. Taking both effects into consideration, we estimate that non-saturated NMD reduces the fluorescence of our TCRβ-GFP fusion protein expressed from the NMD reporter construct ~4- to 6-fold.

**Establishing cell lines stably expressing the NMD reporter**

After having confirmed that the GFP signal proportionally reflects the mRNA level of the NMD reporter, we transfected HT1080 (data not shown) and HeLa cells with our NMD reporter plasmid and selected a number of single cell clones that had integrated the reporter gene into the genome. In order to select a suitable NMD reporter system, we first screened the monoclonal cell lines by real-time RT–PCR for lines expressing a clearly detectable level of the HA-TCRβ-GFP reporter mRNA, which allowed us to eliminate ‘false positives’ that did not express the reporter gene at all (data not shown). Cell lines expressing the reporter gene were then treated with the translation inhibitor CHX for 3 h, which is a standard way to inhibit NMD (31), and the increase in reporter mRNA levels was quantified by real-time RT–PCR. As a control, we also generated and analysed in parallel stably transfected clonal cell lines expressing the ΔJCin construct. Because this construct lacks an intron downstream of the PTC, its mRNA should not be subjected to NMD. Figure 3 shows the relative mRNA levels of a NMD-reporter-expressing cell line (left panel) and of a cell line expressing the ΔJCin control construct (right panel) after the inhibition of NMD with CHX. The NMD reporter mRNA increased on average 6.5-fold upon CHX treatment, similar to the previously tested PTC-containing TCRβ-minigenes (37), whereas the mRNA level of the ΔJCin control construct increased only slightly, confirming that this is not a substrate for NMD.

**RNAi-mediated depletion of hUpf1 and hSmg6 causes increased mRNA and protein levels of the NMD reporter gene**

Treatment of cells with CHX is an easy and quick way to inhibit NMD, but because it blocks all cellular translation, unspecific effects influencing the level of NMD reporter mRNA could also occur. Therefore, we decided to interfere with NMD specifically by knocking down essential NMD factors using RNAi techniques. To this end, we used the pSUPER plasmid (34), from which RNAi-inducing shRNAs can be expressed. To be able to use such a transfection-based RNAi approach for subsequent biochemical analysis, an almost 100% transfection efficiency or a way to eliminate untransfected cells before analysis is required. Since the first option is not always feasible, we opted for the second and introduced into the pSUPER vector the puromycin resistance gene, which allows elimination of the untransfected cells during the course of the RNAi experiment (see Materials and Methods). The NMD reporter gene-expressing HeLa cell line was transfected with such pSUPERpuro plasmids encoding shRNAs against the two essential human NMD factors hUpf1 (12,13,38) or hSmg6 [also called hEST1A or Smg5/7a (18–20)], and the effect on the reporter mRNA level was analysed (Figure 4). Compared with cells transfected with the empty pSUPERpuro (Mock), mRNA levels of the NMD reporter increased between 6- and 11-fold when hUpf1 was knocked down by RNAi (Figure 4A, left panel, and Figure 4C), whereas the ΔJCin mRNA level in the control cell lines was not affected (Figure 4A, right panel). While RNAi against hUpf1 worked quite efficiently, resulting in a 10-fold or greater reduction of hUpf1 mRNA (Figure 4C, and data not shown) and barely detectable hUpf1 protein levels (Figure 4B) under our experimental conditions, RNAi against hSmg6 resulted only in a ~3-fold reduction of hSmg6 mRNA in our hands (Figure 4C, and data not shown). Yet, this moderate reduction of hSmg6 caused the same increase of the NMD reporter mRNA as knockdown of hUpf1 (Figure 4C). This result suggests that hSmg6 might be a limiting factor for NMD function. Collectively, these results show that the expression of our NMD reporter gene increases as predicted when NMD deficiency is induced by RNAi.

**Detection of NMD deficiency by flow cytometry**

In order to use the NMD reporter system as a tool for screening large number of cells for NMD deficiency, its readout (i.e. GFP fluorescence) should be detectable by techniques capable of analyzing many cells within a short time. To detect and isolate NMD-deficient cells in a large cell population, flow cytometry is today’s method of choice. To test the suitability of our NMD reporter system for analysis by flow cytometry, we knocked down hUpf1 and hSmg6 in the reporter gene-expressing HeLa cell line as in the previous experiment and determined the fluorescence emitted by these cells using FACScan. As a reference signal for GFP in NMD-competent cells, the reporter cell line was transfected with the empty pSUPERpuro (Mock), and to control for autofluorescence of the cells, mock-transfected parental HeLa cells were also analysed. The two cell populations with RNAi-mediated NMD deficiency exhibited clearly more GFP fluorescence than the mock-transfected reporter cell line, demonstrating that flow cytometry can be used to analyse the NMD reporter system (Figure 5A). In the mock-transfected reporter cells, almost no GFP signal could be detected over the background caused by the autofluorescence, similar to the situation when small amounts of NMD reporter gene were transiently expressed (Figure 2A). Since almost no GFP signal over background could be detected in mock-transfected reporter cells, the
A

Mock transf.  0.06 %
GFP pos.  MFI = 9.13

50 ng plasmid  0.22 %
GFP pos.  MFI = 8.33

100 ng plasmid  0.65 %
GFP pos.  MFI = 11.83

200 ng plasmid  2.81 %
GFP pos.  MFI = 13.92

400 ng plasmid  11.2 %
GFP pos.  MFI = 17.04

800 ng plasmid  29.4 %
GFP pos.  MFI = 25.63

B  GFP signal

C  mRNA level

D  GFP signal PTC+ versus ΔJCin

| ng transf. plasmid DNA | GFP signal PTC+ | GFP signal ΔJCin | PTC+ / ΔJCin |
|------------------------|----------------|-----------------|-------------|
| 50                     | 1.83           | 16.55           | 0.11        |
| 100                    | 7.69           | 41.02           | 0.19        |
| 200                    | 39.12          | 222.92          | 0.17        |
| 400                    | 190.85         | 641.89          | 0.29        |
| 800                    | 752.50         | 2074.31         | 0.36        |
| none                   | 0.55           |                 |             |
fold increase upon knockdown of hUpf1 and hSmg6 cannot be calculated. As a control, NMD reporter mRNA levels were measured from the same cells analysed by flow cytometry. As in previous experiments (Figure 4C), the RNAi-mediated knockdown of hUpf1 and hSmg6 resulted in an ~10-fold increase of reporter mRNA (Figure 5B). RNA from the mock-transfected parental HeLa cells gave no detectable signal in the real-time RT–PCR assay, confirming that the TaqMan probe for measuring reporter mRNA is specific (data not shown). In summary, this experiment shows that our NMD reporter system can be used in conjunction with flow cytometry to detect NMD-deficient cells.

Detection of NMD deficiency using a microplate spectrofluorometer

For other types of automated large-scale screens, the preferred analysis method includes the use of a fluorescence plate reader. A typical example for such an application would be a screening of chemical libraries for compounds that inhibit NMD. Although the nature of the reporter system would prevent the identification of compounds that inhibit NMD by blocking translation, it might still be very useful to identify novel compounds that inhibit NMD through other mechanisms, such as binding to essential NMD factors. We, therefore, wanted to test whether our NMD reporter cells can be used to detect NMD deficiency by measuring the GFP fluorescence in whole cell extracts using a microplate spectrofluorometer (SpectraMAX Gemini). To this end, hUpf1 was again knocked down in HeLa cell lines stably expressing the HA-TCRβ-GFP reporter gene together with a constant amount of a β-globin-expressing plasmid (41). The HeLa cells were analysed by flow cytometry. The gate for GFP positive cells was set so that the autofluorescence of the mock-transfected cells did not score as GFP positive. MFI, mean fluorescence intensity. (B) MFI values multiplied with the percentage of GFP positive cells from (A) were plotted against the amount of transfected NMD reporter plasmid DNA. (C) From an aliquot of the cells used in (A), RNA was isolated and relative NMD reporter mRNA and β-globin mRNA was measured by real-time RT–PCR. Relative reporter mRNA levels were normalized to relative β-globin mRNA levels, and average values and standard deviations of three real-time PCR runs are shown. (D) Analogous to the experiment shown in (A) and (B) for the PTC+ reporter gene, the GFP signals from cells transiently transfected with different amounts of the ΔJCin control construct were determined by flow cytometry. As in (B), the GFP signal is defined as MFI* % of GFP positive cells.
down in the NMD reporter cell line, and mock-transfected reporter cells and mock-transfected plain HeLa cells were used as controls. Whole cell extracts of each cell population were prepared and the fluorescence at 512 nm (GFP) of different dilutions thereof was measured. The values obtained with the NMD reporter cells were normalized by subtracting the background values measured with plain HeLa (Figure 6A). Very similar to the result of the flow cytometry analysis, no GFP signal significantly over background could be detected in the NMD-competent, mock-transfected reporter cells where hUpf1 was knocked down. This demonstrates that...
in general the NMD reporter system can also be used in combination with a fluorescence plate reader. However, the minimal amount of lysate required to detect a GFP signal clearly over background in the hUPF1-depleted cells corresponds to ~4 × 10⁵ cells, which is too much for a large-scale screening. Thus, in order to use the NMD reporter system for spectrofluorometer-based drug discovery, the sensitivity of the assay needs to be improved (see below).

To assess the efficiency of the hUpf1 knockdown and its effect on the level of reporter protein, aliquots of the cells used in Figure 6A were analysed by western blot analysis (Figure 6B). Probing with the anti-hUPF1 antibody shows that in this experiment too, the expression of shRNAs targeting hUpf1 mRNA led to a strong reduction in hUPF1 protein. With the anti-HA antibody, NMD reporter protein could be detected in the hUpf1 knockdown cells, but not in the mock-transfected cells, consistent with result from the spectrofluorometer. The monoclonal antibody Y12, which detects the Sm proteins B, B', D1 and D3 (39), was used as a loading control. Real-time RT–PCR showed that the NMD reporter mRNA levels increased 3.2-fold in this particular experiment (data not shown).

**Detection of NMD-deficiency by fluorescence microscopy**

The third detection method that we tested on our NMD reporter cells was fluorescence microscopy using a confocal laser scanning microscope. As previously, hUpf1 was knocked down in the NMD reporter-expressing HeLa cell line by transfection of shRNA targeting pSUPERpuro following by selection of the puromycin-resistant cells. Four days after transfection, these cells were examined under the fluorescence microscope and compared with the mock-transfected, NMD-competent reporter cells and untransfected HeLa cells. Only weak green fluorescence could be detected mainly in the cytoplasm of mock-transfected reporter cells when the pinhole of the confocal microscope was opened to achieve maximal sensitivity (Figure 7D). The average fluorescent signal was nevertheless significantly higher than the autofluorescence seen in untransfected HeLa cells (Figure 7B), and we therefore conclude that in contrast to flow cytometry (Figure 5A), the low basal GFP level of the NMD reporter can be detected by laser scanning fluorescence microscopy. In contrast to this low basal level of GFP in the mock-transfected reporter cells, the green fluorescence, which is predominantly cytoplasmic, is much more intense of hUPF1-depleted reporter cells (compare Figure 7D and F). Differential interference contrast (DIC) images of the same cells showed no obvious morphological changes 4 days after transfection of pSUPERpuro plasmids (compare Figure 7C and E with Figure 7A), suggesting that at this time point, abrogation of NMD has not yet severely damaged the cells (see below). We conclude that our NMD reporter system can also be used to screen for individual cells by fluorescence microscopy.

**CONCLUSIONS**

Here, we describe the construction of a GFP-based reporter system, in which increased green fluorescence of the cells signals NMD deficiency. We validate this reporter system by demonstrating that increased GFP signals are in fact detected by flow cytometry, spectrofluorometry and fluorescence microscopy in cells, where NMD has been abrogated by knocking down essential NMD factors using RNAi techniques. In all these tests, our NMD reporter system behaved as expected, proving its potential to become a valuable tool for future studies of NMD.

If the GFP-based NMD reporter system is used to screen for NMD-deficient cells, one must be aware that, for example, mutants that enhance transcription or translation, or that inhibit general mRNA turnover or protein degradation will also result in increased GFP fluorescence. To identify such false positives, samples with increased GFP signal must therefore be validated by further experiments. To this end, reporter mRNA levels and mRNA of an endogenous gene can be measured, and in addition, expression and analysis of another well-established NMD reporter mRNA can be used to confirm NMD deficiency.

For some automated large-scale screenings, the currently generally low GFP signal emitted by the reporter may limit its application. However, there are several ways to improve the sensitivity of the system if required. On the one hand, detection systems are becoming more and more sensitive, and on the other hand, it should be possible to generate cell lines that contain more copies of the HA-TCRβ-GFP gene and, thus, would be expected to generate more fluorescence per cell. For flow cytometry applications, substitution of GFP in our reporter gene by another fluorescent protein variant that emits light at a different wavelength might overcome the current problem of low GFP signals being hidden in the autofluorescence of the cells (Figures 2A and 5A). To increase the sensitivity of the reporter system in spectrofluorometric applications, GFP could be replaced by firefly luciferase. Because luciferase produces chemiluminescence in an enzymatic reaction by oxidation of luciferin, detection of more light per reporter molecule would be expected when using a microplate spectrofluorometer.

In parallel to the HeLa NMD reporter system mainly described here, we have also generated a diploid HT1080 cell line that expresses our NMD reporter gene (Figure 1C, and data not shown) and used these cells for a mutagenesis screen with the goal of obtaining NMD-deficient cell lines. The mutagenesis scheme and the selection of high GFP-expressing cells by flow cytometry was essentially as described for a similar screen, which yielded mutant cell clones deficient for rapid ARE-dependent IL-3 mRNA turnover (26). Although high GFP-expressing cells occurred at a very low frequency in our mutated cell population, we were never able to proliferate the high GFP-expressing cells after sorting (data not shown). This was not due to cell culturing problems, because from the mutated cell pool, cell lines from single, low GFP-expressing cells could be obtained with a reasonable plating efficiency (data not shown). Meanwhile, we believe that the reason for the failure of this mutagenesis screen might be that NMD could be essential for survival of mammalian cells, even in cell culture. In our RNAi experiments targeting hUpf1 and hSmg6, the cells die 7–8 days after transfection (A. Paillusson and O. Mühlemann, unpublished data). In addition, from all attempts to stably transfect pSUPERpuro targeting hUpf1, we only obtained cell clones in which the hUpf1 protein level was reduced 5-fold or less (data not shown), while hUpf1 is usually reduced ~10-fold.
or more when cells transfected with the same pSUPERpuro plasmids are assayed 4 or 5 days after transfection (Figures 4B and 6B, and data not shown). Furthermore, Upf1 knock-out is lethal in mice, with embryos dying 6–7 days post coitus (40). All these observations are consistent with the suggestion that NMD might be essential for viability of mammalian cells. However, it is also possible that hUpf1 and hSmg6 play additional roles in other essential cellular processes.

Figure 7. Detection of NMD deficiency by confocal laser scanning fluorescence microscopy. The NMD reporter cell line was grown on coverslips and transfected with the empty pSUPERpuro plasmid (mock; C and D) or with pSUPERpuro-Upf1 (E and F) as in Figure 4. Untransfected HeLa cells served as a control for background autofluorescence (A and B). After fixation, the cells were analysed using a confocal laser scanning fluorescence microscope. Differential interference contrast (DIC) images (A, C and E) and fluorescence images in the GFP channel (B, D and E) of the same cells are shown.
In conclusion, the GFP-based NMD reporter gene system described here faithfully reports NMD deficiency in individual cells by increasing the fluorescence of these cells. Because the NMD reporter system functions in combination with flow cytometry, spectrophotometry and fluorescence microscopy, it has the potential to be used for automated large-scale screening applications, although some modifications might be necessary for specific applications (see above). With these properties, this NMD reporter gene system represents a new and valuable tool in the toolbox of researchers investigating the molecular mechanisms of mammalian NMD.

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