Translation termination is signaled by three stop codons: UAA, UAG, and UGA. This mechanism is highly conserved, although each stop codon has a different efficiency for terminating translation. UGA is considered to be a “leaky” stop codon with the highest intrinsic readthrough potential. UAA shows high fidelity and little intrinsic readthrough potential, whereas UAG has intermediate fidelity (Weiner and Weber, 1973; Lovett et al., 1991). Nonsense mutations create primary premature termination codons (PTCs) and result in either no formation of the target protein or truncated protein with impaired stability.

It is estimated that 30% of human disease-causing alleles are nonsense mutations (Mendell and Dietz, 2001). Other types of mutation, such as frameshift and splicing mutations, lead to secondary PTCs; however, these are not therapeutic targets for readthrough compounds (RTCs). Considering that >1,800 distinct genetic disorders are caused by nonsense mutations, the readthrough of primary PTCs has treatment potential for large numbers of patients.

To date, most reported PTC-RTCs that are active in mammalian cells have belonged to the aminoglycoside antibiotics class (Keeling and Bedwell, 2005; Zingman et al., 2007). Certain types of aminoglycosides can induce ribosomes to read through PTC mutations via insertion of nonaminoglycoside compounds induce readthrough of nonsense mutations

Liutao Du,1 Robert Damoiseaux,5 Shareef Nahas,1 Kun Gao,2 Hailiang Hu,1 Julianne M. Pollard,1 Jimena Goldstine,3 Michael E. Jung,6 Susanne M. Henning,2 Carmen Bertoni,4 and Richard A. Gatti1,3

1Department of Pathology and Laboratory Medicine, 2Center for Human Nutrition, 3Department of Human Genetics, and 4Department of Neurology, David Geffen School of Medicine, 5Molecular Shared Screening Resources, California NanoSystems Institute, 6Department of Chemistry and Biochemistry, University of California, Los Angeles, Los Angeles, CA 90095

Large numbers of genetic disorders are caused by nonsense mutations for which compound-induced readthrough of premature termination codons (PTCs) might be exploited as a potential treatment strategy. We have successfully developed a sensitive and quantitative high-throughput screening (HTS) assay, protein transcription/translation (PTT)–enzyme-linked immunosorbent assay (ELISA), for identifying novel PTC-readthrough compounds using ataxia–telangiectasia (A-T) as a genetic disease model. This HTS PTT–ELISA assay is based on a coupled PTT that uses plasmid templates containing prototypic A-T mutated (ATM) mutations for HTS. The assay is luciferase independent. We screened ~34,000 compounds and identified 12 low-molecular-mass nonaminoglycosides with potential PTC-readthrough activity. From these, two leading compounds consistently induced functional ATM protein in ATM-deficient cells containing disease-causing nonsense mutations, as demonstrated by direct measurement of ATM protein, restored ATM kinase activity, and colony survival assays for cellular radiosensitivity. The two compounds also demonstrated readthrough activity in mdx mouse myotube cells carrying a nonsense mutation and induced significant amounts of dystrophin protein.

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a random amino acid by near-cognate transfer RNA. The therapeutic potential of aminoglycosides has been evaluated in the laboratory for different genetic models, such as cystic fibrosis (Howard et al., 1996; Bedwell et al., 1997; Du et al., 2002), muscular dystrophy (Howard et al., 2000; Wagner et al., 2001; Dunant et al., 2003; Loufrani et al., 2004), Hurler syndrome (Keeling et al., 2001), cystinosis (Helip-Wooley et al., 2002), spinal muscular atrophy (Sossi et al., 2001), ataxia-telangiectasia (A-T; Lai et al., 2004), and type 1 Usher syndrome (Rebibo-Sabbah et al., 2007). Clinical trials also indicate that aminoglycosides can induce some functional protein production; however, the therapeutic benefits remain uncertain (Wilschanski et al., 2000; Clancy et al., 2001; Wagner et al., 2001; Polito et al., 2003). Furthermore, the toxicity of most commercial aminoglycosides in mammals has greatly diminished their potential for successful readthrough therapy (Mingeot-Leclercq and Tulkens, 1999; Guan et al., 2000). Therefore, efforts are underway to develop better aminoglycoside derivatives with reduced toxicity and enhanced activity (Nudelman et al., 2006; Rebibo-Sabbah et al., 2007). Recently, PTC Therapeutics (South Plainfield, NJ) described a more efficient nonaminoglycoside RTC, PTC124, which was developed synthetically by screening >800,000 chemicals and analogues using a luciferase-based high-throughput screening (HTS) assay (Welch et al., 2007; Hirawat et al., 2007; M. Du et al., 2008). A phase-I clinical study in cystic fibrosis confirmed that PTC124 is generally well tolerated and appears to have more efficient readthrough activity than aminoglycosides (Hirawat et al., 2007). Moreover, PTC124 does not induce ribosomal readthrough of normal stop codons. A phase-II clinical trial is underway (Kerem et al., 2008). However, a recent study indicates that the initial discovery of PTC124 by HTS may have been biased by its direct effect on the FLuc (firefly luciferase) reporter used (Auld et al., 2009), indicating the importance of a luciferase-independent HTS assay for future drug screening.

In an effort to discover new RTCs, we developed a sensitive and quantitative luciferase-independent HTS assay, protein transcription/translation (PTT)–ELISA. The PTT-ELISA assay was validated for a fully automated 384-well robotic platform and used to screen ~34,000 compounds. We focused followup efforts on 12 low-molecular-mass nonaminoglycoside compounds. From there, we identified two compounds that induced low levels of full-length functional A-T mutated (ATM) protein in A-T cells carrying ATM nonsense mutations, as demonstrated by direct measurement of ATM protein using ATM-ELISA, ATM-Ser1981 auto-phosphorylation, trans-phosphorylation of structural maintenance of chromosome (SMC) 1–Ser966, and colony survival assay (CSA). Both compounds also showed PTC-readthrough activity in mdx mouse myotube cells carrying a nonsense mutation and induced significant amounts of dystrophin protein.

Collectively, these studies provide the first robust luciferase-independent HTS assay for identifying RTCs and proof of principle for PTC readthrough by nonaminoglycoside compounds. They further establish that enhanced PTC readthrough can be considered as a therapeutic strategy for correcting nonsense mutations in many genetic diseases.

RESULTS
Development and validation of HTS assay PTT-ELISA
In vitro PTT was originally designed to detect truncating mutations (Roest et al., 1993; Telatar et al., 1996). ELISA was coupled into PTT to improve throughput for the detection of these mutations (Gite et al., 2003; L. Du et al., 2008). In a previous study, we used a PTT–PAGE gel approach to evaluate various aminoglycosides for readthrough activity of ATM nonsense mutations (Lai et al., 2004). However, the gel-based PTT assay was time consuming and involved the use of radioactive material; thus, it was difficult to automate for a high-throughput format. Herein, we have developed a plasmid-driven PTT-ELISA assay for screening large numbers of compounds for PTC readthrough activity. The assay uses plasmid templates containing prototypic ATM mutations, patterned after specific disease-causing ATM mutations. So as to work in a mammalian system, rabbit reticulocytes were chosen to drive the PTT reaction. Various fragments of mutated ATM alleles from cells of A-T patients were cloned into the plasmids and were N- and C-terminally tagged with the epitopes myc and V5, respectively. Anti-myc antibody was used to capture the translated protein onto an ELISA plate. If compounds induce PTC-readthrough in the assay, the plasmid-driven PTT results in a full-length ATM fragment including the V5 tag, which is detected with anti–V5–horseradish peroxidase (HRP) antibody (Fig. 1 a).

We constructed three mutant plasmids that contain prototypic PTC mutations in the ATM gene from three different A-T patients. The plasmid used for preliminary screening, plasmid–TAT51, contains ATM region 5 fragment (codons 1403–1886) and harbors a nonsense (PTC) mutation (c.5623C→T) that leads to a TGA C stop codon. The second mutant plasmid, plasmid–AT153LA, contains the same TGA stop codon but at a different position within the gene (c.8977C→T) in region 8 (codons 2550–3050) and also has a different +4 nt (TGA A). This was used to monitor the effect of surrounding sequences of PTCs on readthrough ability. A third mutant plasmid, plasmid–AT185LA, contains a different stop codon, TAA G, resulting from a nonsense mutation (c.3673C→T) in region 4 (codons 1041–1531). Plasmids containing the same fragments but without mutations were constructed by in vitro mutagenesis of patient-derived complementary DNA and were used as wild-type controls.

We first evaluated the specificity and sensitivity of PTT-ELISA to detect the in vitro–translated full-length protein fragment. All three mutant plasmids gave only background signal, whereas the comparable wild-type plasmids showed signals greater than ~200-fold over background (Fig. 1 b, top), indicating that the assay specifically recognized full-length protein fragments. We next used TAT51 wild-type plasmid to evaluate the sensitivity of the assay. The TAT51 wild-type plasmid was serially diluted with TAT51 mutant plasmid and...
used to drive PTT reactions. The sample containing 1.2% of wild-type plasmid (2/158 ng) still gave signal that was two-fold higher than that of mutant plasmid (Fig. 1 b, bottom), establishing the sensitivity of the assay at ~1%. The sensitivity of an HTS assay is especially important for RTC screening because all RTCs to date have been only weak PTC readthrough inducers, and new classes of RTCs are expected only to be identified with a highly sensitive screening assay.

Next, we used two well known RTCs, G418 and gentamicin, to test the efficiency of the PTT-ELISA assay. Both compounds showed significant PTC readthrough activity with TAT51 mutant plasmid (TGA C) over a large dose range (40 nM–10 µM; Fig. 1 c). Moreover, G418 showed obvious toxicity for the PTT reaction at concentrations >2.5 µM, whereas gentamicin toxicity was not observed until 12.5 µM. These results are consistent with our previously published S35-PTT gel data (Lai et al., 2004) and demonstrate that the PTT-ELISA readily identifies compounds with readthrough activity.

We next validated the assay for use in a fully automated 384-well robotic platform, using plasmid-TAT51 (TGA C) as readthrough template and 1 µM G418 as the positive readthrough control. To further reduce the cost of the assay, we optimized a decrease in the PTT reaction volume from 25 to 5 µl. As shown in Fig. 1 d, the wells containing G418

Figure 1. HTS PTT-ELISA assay for RTCs screening. (a) Schematic of PTT-ELISA HTS. ATM regions containing a PTC mutation were cloned into plasmids and tagged with c-myc and V5 epitopes at each end. Compounds with PTC readthrough activity induce full-length protein, which can be identified by anti–V5-HRP. (b) Specificity and sensitivity of PTT-ELISA. All three mutant plasmids showed only background readings, whereas wild-type plasmids showed signals >200-fold over background (P < 0.0001), indicating that the assay specifically detects full-length proteins. The sample containing 2 ng of wild-type plasmid still gave signals that were twofold higher than those of mutant plasmid alone (P < 0.01), indicating that the sensitivity of the assay is ~1% (2/158 ng). Error bars indicate the variation of duplicate samples. (c) Assay evaluation using aminoglycosides. Both G418 and gentamicin induced significant PTC readthrough over a large dose range. (d) Readings for a sample row from a 384-well plate. Samples containing G418 exhibited significantly different signal (Z’ factor > 0.6) over the no-drug wells (P < 0.0001). ****, P < 0.0001, as compared with control. Experiments were repeated three times for b and c and five times for d.
exhibited significantly different signals (Z’ factor > 0.8) over the no-drug wells. The fully automated assay also showed consistent plate-to-plate accuracy and efficiency, indicating the suitability of the assay for HTS (unpublished data).

Screening of chemical libraries for RTCs
About 34,000 compounds were screened to discover novel RTCs. Plasmid-TAT51 (TGA C) was used for the initial screening. Each compound was screened at a final concentration of 10 µM in the assay mixture. For each screening plate, we included both positive (with 1 µM G418) and negative (with DMSO) samples as quality controls. Samples with signal >2-fold over the negative control were scored as potential “hits.” Our initial screens yielded 12 low-molecular-mass RTCs with appreciable readthrough activity. All 12 hits were then confirmed by manual PTT-ELISA at multiple concentrations (chemical names shown in Table S1). None of them were aminoglycosides and none of them had been previously reported. The chemical names and structures of the two leading compounds, RTC#13 and #14, are shown in Fig. 2 a; the names and structures for the remaining 10 RTCs are given in Fig. S1 and Table S1.

The EC_{50} of five compounds (RTC#4, #11, #13, #14, and #16) were <10 µM (Fig. 2 b, RTC#13 and #14), implying therapeutic potential of those compounds. Notably, in these initial cell-free experiments, the maximum in vitro readthrough effect for the new compounds was not as favorable as that of G418 or gentamicin. For example, the maximum readthrough activity of RTC#13 and #14, detected by the cell-free PTT-ELISA assay, was ~10% of the maximum activity of G418 and gentamicin in the same assay (Fig. 1 c vs. Fig. 2 b). This may be related to the solubility, permeability, or toxicity of the compounds. In contrast, both RTC#13 and #14 had an EC_{50} <10 µM, they were less toxic, and they did not show obvious inhibition of PTT at high concentrations (>50 µM), unlike both G418 and gentamicin (Fig. 1 c).

The readthrough efficiency of these compounds was also tested on two other stop codon contexts, TGA A and TAA G. For TGA A, all 12 compounds showed various extents of readthrough activity (unpublished data). For TAA G, six compounds (RTC#10, #11, #13, #14, #16, and #17) showed appreciable activity (unpublished data). The data for RTC#13 and #14 are shown in Fig. 2 c.

RTC-induced ATM protein and ATM-Ser1981 phosphorylation in A-T lymphoblastoid cell lines (LCLs)
To test whether the newly identified compounds could induce ATM protein in A-T cells with PTC mutations, we exposed A-T LCLs to each compound for 4 d before harvesting the cells. Western blotting of nuclear lysates was generally of insufficient sensitivity for monitoring RTC-induced ATM protein (Lai et al., 2004); therefore, we used an ATM-ELISA method to measure intranuclear ATM protein in cells (Butch et al., 2004). Concentrations of individual RTCs used to treat AT153LA LCL with a homozygous TGA A mutation were based on their cytotoxicity profile (Fig. S2, RTC#13 and #14). Cells treated with the highest RTC concentrations still had >70% viability as compared with untreated cells. RTC#13 and #14 consistently induced low but detectable levels of ATM protein in A-T LCLs (Fig. 3 a). The restored ATM protein was ≤5% of the wild-type LCLs. Gentamycin and G418 also induced small amounts of ATM protein (Fig. 3 a). We were encouraged by these results because A-T patients with some residual ATM protein levels can have substantial ATM kinase activity, and this has been sometimes associated with a later onset and slower progression of symptoms (Gilad et al., 1998; Chun et al., 2003). Thus, we believe that even modest increases in ATM protein levels have therapeutic potential.

To test whether the newly identified compounds could induce ATM-Ser1981 phosphorylation in A-T lymphoblastoid cell lines (LCLs), we exposed A-T cell lines to each compound for 4 d before harvesting the cells. Our previous studies had shown that, at this concentration, G418 induced a maximum level of ATM-Ser1981 IRIF (Lai et al., 2004). As anticipated, significant IRIFs were induced in G418-treated cells. The level in A-T cells was about half that in wild-type cells. The compounds were also tested in another A-T cell line, AT229LA, with homozygous TAG A mutations, and both RTC#13 and #14 induced significant numbers of ATM-Ser1981 IRIFs in A-T cells with TAG A, whereas only a background level of IRIFs was observed in the untreated controls (Fig. 3 b). At same time, ~51% of wild-type cells showed distinct post-IR foci. The maximum IRIF induction achieved in A-T cells by 10 µM RTC#13 was ~40% of wild-type level. As a positive readthrough control, 144 µM of G418 (100 µg/ml) was used to treat cells. Our previous studies had shown that, at this concentration, G418 induced a maximum level of ATM-Ser1981 IRIF (Lai et al., 2004). As anticipated, significant IRIFs were induced in G418-treated cells. The level in A-T cells was about half that in wild-type cells. The compounds were also tested in another A-T cell line, AT229LA, with homozygous TAG A mutations, and both RTC#13 and #14 induced significant numbers of ATM-Ser1981 IRIF, as compared with non-treated A-T cells (Fig. 3 c).

To further compare the readthrough activity of RTC#13 and #14 with gentamicin and G418, we used a flow cytometry (FC)–based ATM-Ser1981 phosphorylation assay to assess readthrough activity in AT229LA cells (TAG A; Fig. 4). Both RTC#13 and #14 induced increased ATM-Ser1981 auto-phosphorylation, as indicated by the right shift of fluorescence intensity (FI). This was consistent with previous ATM IRIF data (Fig. 3 c). Encouragingly, ATM-Ser1981 phosphorylation levels induced by RTC#13 and #14 were similar to those induced by the same concentrations of gentamicin and G418 in the same cell line (Fig. 4). Neither of the two compounds produces green autofluorescence (isotype controls; Fig. 4, top right histogram). Similar readthrough effects were also observed in AT153LA cells (TGA A) using cytometry-based ATM-Ser1981 phosphorylation assay (unpublished data).

RTC-induced restoration of SMC1-Ser966 phosphorylation in A-T LCLs
As an alternative assay to assess ATM kinase function, we used a recently developed FC–based assay to measure the trans-phosphorylation of SMC1 in A-T LCLs (Fig. 5; Nahas et al., 2009). ATM phosphorylates SMC1 at Ser966 and Ser957...
after IR-induced double-strand breaks (Yazdi et al., 2002; Kitagawa et al., 2004), and IR-induced SMC1 phosphorylation is deficient in A-T cells. In untreated AT153LA cells (TGA A), we observed no phosphorylation of SMC1 (Fig. 5 a, top, left). After treatment with either RTC#13 or #14, a right shift in FI was observed, indicating restored ATM kinase activity. Both G418 and gentamicin induced SMC1 phosphorylation in A-T cells at similar levels. We also tested the same compounds on TAG A mutation using AT229LA cells. They all induced detectable SMC1 phosphorylation (Fig. 5 b).

RTC-induced correction of SMC1 phosphorylation was dose dependent in AT153LA cells (TGA A; Fig. S3).

RTC-induced restoration of ATM kinase in A-T fibroblast cells
To determine whether the two lead RTCs were also active in other types of cells, we tested them against the A-T fibroblast cell line GM02052, which contains a homozygous TGA G mutation (c. 103C→T). RTC-treated cells showed slightly increased IR-induced SMC1-Ser966 (Fig. 6 a) and ATM-Ser1981 phosphorylation (Fig. 6 b), as compared with untreated

Figure 2. Two leading RTCs were identified by HTS. (a) Molecular structures of RTC#13 and #14 (2-imino-5-[(5-(2-nitrophenyl)-2-furylmethylene)-1,3-thiazolidin-4-one and 4-tert-butyl-2-[(3-nitrobenzylidene)amino]pheno). (b) TGA A readthrough activity by PTT-ELISA. (c) TGA A and TAA G readthrough activity. **, P < 0.01; ***, P < 0.001, as compared with untreated control. Experiments were repeated four times for b and three times for c.
Because the radiosensitivity of A-T cells results from the ATM deficiency, we next investigated whether RTC#13 and #14 can abrogate A-T cell's radiosensitivity of A-T cells.

RTC-induced cell survival correction in A-T LCLs

Figure 3. RTCs induced intranuclear ATM protein and post-IR ATM-Ser1981 foci in A-T LCLs. LCLs were treated with compound for 4 d before harvesting. Induced ATM protein was assessed by ATM-ELISA and ATM-Ser1981 IRIF. For IRIF test, cells were irradiated with 2 Gy and IRIFs were scored after 30 min. All experiments were repeated three times. (a) Cells treated with various doses of compound RTC#13 and #14 showed a significantly increased ATM protein level, as compared with nontreated A-T samples (P < 0.05). The dashed line indicates the basal ATM protein level in untreated A-T cells. (b) RTC-induced ATM-Ser1981 IRIF in AT153LA (TGA A) cells. (c) RTC-induced ATM-Ser1981 IRIF in AT229LA (TAG A) cells. *, P < 0.05; **, P < 0.01, as compared with untreated sample. Error bars indicate the variation of two independent experiments.
Figure 4. RTCs induced FC-ATM-Ser1981 phosphorylation in A-T LCLs. AT229LA cells (TAG A) were treated with RTC#13 and #14 for 4 d and analyzed for ATM-Ser1981 phosphorylation using FC. Gentamicin and G418 were used as positive readthrough controls. All compounds induced ATM phosphorylation in A-T cells, as indicated by a right Fl shift. Neither compound produced autofluorescence, as shown in the top right histogram. Results were consistent in three independent experiments.
AT153LA cells (TGA A) were treated with compounds for 4 d and followed by CSA. One wild-type cell line and one different A-T cell line were used as assay quality controls. As expected, the wild-type and A-T controls showed 49 and 11% survival fraction, respectively. These were within the normal (>36%) and radiosensitive (<21%) range, respectively (Fig. 7, right two bars). Untreated AT153LA cells showed a 13.5% survival fraction, which is radiosensitive (Fig. 7, left bar). Encouragingly, we found that both RTC#13 and #14 (at 10 and 20 µM) abrogated AT153LA cell radiosensitivity, from radiosensitive (13.5%) to intermediate (21–36%) range. 100 µM G418 also abrogated radiosensitivity to intermediate range, whereas gentamicin did not show a significant effect in this assay at the tested concentrations (10 and 20 µM). The definition of CSA range used was previously established in our laboratory and is used for clinical A-T diagnosis (see Materials and methods).

**RTCs induced *mdx* PTC readthrough in mouse myotube cells**

To investigate the ability of RTC#13 and #14 to readthrough a PTC mutation in genes other than *ATM*, we used *mdx* myotubes. The *mdx* mouse has been widely used as a model for Duchenne muscular dystrophy. It carries a C to T transition in exon 23 of the dystrophin gene that creates a premature stop codon (TAA), resulting in an absence of dystrophin protein (Sicinski, et al., 1989). Because gentamicin has been shown to induce readthrough of the *mdx* PTC mutation and to restore dystrophin expression in *mdx* mice both in vitro and in vivo (Barton-Davis, et al., 1999), we used it as a positive readthrough control in the experiments. Muscle progenitor cells isolated from *mdx* mice were induced to differentiate for 24 h before addition of gentamicin or compounds RTC#13 and #14. Myotube cultures were exposed to compounds for 72 h and then analyzed for dystrophin by Western blotting. Full-length dystrophin was clearly detected in cells treated with RTC#13 and #14 at concentrations of 10 and 20 µM but not in cultures treated with gentamicin at concentrations of 20 µM (unpublished data). We next used immunoprecipitation to increase the sensitivity of the assay (Fig. 8). Dystrophin expression was present in cells treated with gentamicin at concentrations of 20 and 100 µM. Both of our lead compounds showed readthrough activity, inducing significant

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**Figure 5. RTCs induced FC-SMC1 pSer966 phosphorylation in A-T LCLs.** (a) RTC#13 and #14 restored SMC Ser966 phosphorylation in AT153LA cells [TGA A]. (b) RTC#13 and #14 restored SMC1 Ser966 phosphorylation in AT229LA cells [TAG A]. All experiments were repeated three times.
amount of dystrophin protein. Results were consistent in multiple experiments performed.

**DISCUSSION**

We believe that HTS can identify new nonaminoglycoside RTCs with therapeutic potential. In this study, we used A-T as a disease model to identify new RTCs. A-T is a progressive autosomal recessive neurodegenerative disorder resulting from mutations in the *ATM* gene (Perlman et al., 2003; Chun and Gatti, 2004). ATM protein plays a very important role in cell cycle control, DNA damage repair, the oxidative stress response, and apoptosis (Shiloh, 2006). The A-T disorder provides an appropriate laboratory model for demonstrating novel principles of mutation-targeted therapy. In the *ATM* mutation spectrum, primary nonsense mutations account for \( \sim 15\% \) of the unique mutations detected in A-T patients (www.lovda.nl/ATM). A well characterized spectrum of *ATM* mutations, supported by an extensive library of LCLs derived from patients with those mutations, allowed us to investigate the effect of nonaminoglycoside RTCs on various primary premature stop codons. We have previously used aminoglycosides and antisense oligonucleotides to correct nonsense and splicing *ATM* mutations, respectively, and to restore functional ATM protein in A-T LCLs (Lai et al., 2004; Du et al., 2007). These studies suggest that therapeutic benefits might be achieved if even modest increases in functional ATM protein levels can be induced. In these limited studies, we saw no significant effects on efficiency of readthrough from the fourth nucleotide of each tested stop codon.

Herein, we successfully developed a sensitive luciferase-independent HTS assay by coupling PTT and ELISA. PTT-ELISA shows high specificity for detecting readthrough products and, thus, minimizes false positives in the initial large-scale library screening. The assay is also very sensitive; the minimum detection threshold is \( \sim 1\% \), which ensures its efficiency as a HTS assay. The efficiency of PTT-ELISA was further evaluated using two well known RTCs, G418 and gentamicin. The assay was able to detect their readthrough amount of dystrophin protein. Results were consistent in multiple experiments performed.

**Figure 6.** RTCs restored ATM kinase activity in A-T fibroblast cells. GM02052 cells with a homozygous c.103C→T mutation (TGA G) were treated with RTC#13 and #14 for 4 d, and ATM kinase activity was assessed using FC-based SMC1-Ser966 phosphorylation and ATM-Ser1981 phosphorylation. All experiments were repeated three times. (a) Histograms of FC-SMC1-Ser966. (b) Cell population positive for ATM-Ser1981 staining. The dashed line indicates the basal ATM s1981 phosphorylation in the nontreated A-T cells after radiation (10 Gy). **, P < 0.01, as compared with untreated sample. Error bars indicate the variation of two independent experiments.
activity over a very large concentration range (G418, 40 nM–10 µM; gentamicin, 40 nM–100 µM). Furthermore, PTT-ELISA has been validated for a fully automated robotic platform, with consistent accuracy between plates. The 384-well format dramatically reduced the workload for screening thousands of compounds and also saved time, costs, and reagents. Moreover, this assay has the potential to be validated for a 1,536-well format. For these reasons, we believe that the PTT-ELISA HTS assay provides a powerful new tool for identifying new RTCs.

From a library of ~34,000 compounds, we identified 12 low-molecular-mass compounds (between ~300 and 450 daltons) with PTC readthrough activity. None of these new compounds were aminoglycosides. Several compounds showed EC₅₀ values <10 µM, implying their potential for further development. To further assess these compounds in cell systems, we tested their readthrough activity in A-T LCLs with a variety of nonsense mutations. We used ATM-ELISA to directly detect ATM protein levels in treated cells. Subsequently, more sensitive cell-based assays, such as FC-based SMC1-Ser966, ATM-Ser1981 phosphorylation, and ATM-Ser1981-IRIF, were used to assess restored ATM kinase activity. Among the 12 compounds, RTC#13 and #14 showed PTC readthrough activity in A-T cells, both in LCLs and fibroblasts, as demonstrated by ATM-ELISA, ATM kinase activity (autophosphorylation of ATM and trans-phosphorylation of SMC1), and CSA. To determine whether the compounds can read through PTC mutations in other genes, we selected mouse mdx myotube cells and tested their ability to induce readthrough in a different species, a different cell type (nondividing muscular cells), and a different premature stop codon (TAA). In other studies, the TAA codon has proven the most difficult to read through (Kimura et al., 2005; Welch et al., 2007). Both RTC#13 and #14 induced PTC readthrough of the mouse mdx dystrophin gene.

Collectively, our data showed that both RTC#13 and #14 had comparable readthrough activity in cell-based assays, even though their activities in cell-free PTT-ELISA assay were much lower than gentamicin and G418. This may be related to differences in cell-based metabolism, drug solubility, and permeability of compounds.

To assess the impact of the compounds on readthrough of normal stop codons of other proteins, we performed two-dimensional gel electrophoresis. Neither compound significantly interfered with protein expression patterns (Fig. S4), implying that these compounds have potential for further

**Figure 7.** RTCs abrogated the radiosensitivity of A-T LCLs. AT1S3LA cells (TGA A) were treated with compounds and the CSA was measured. RTC#13, RTC#14, and G418 increased cell survival fractions to intermediate range. Gentamicin did not show an effect at tested concentrations. Results were consistent in three independent experiments.
development. However, we anticipate that therapeutic effects may not be discernable in A-T patients if the RTC-induced ATM level is <15% of normal, and this very likely also applies to other genetic disorders. Therefore, further structural modifications to improve pharmodynamics will be necessary. Structural optimization has improved the readthrough activity and lowered the toxicity of aminoglycosides (Nudelman et al., 2006; Rebibo-Sabbah et al., 2007). Therefore, it is expected to be effective for developing better nonaminoglycoside readthrough analogues as well.

The underlying mechanisms of PTC readthrough activity for these newly identified compounds remain unknown. It has been demonstrated that all of the known PTC-RTCs function by interfering with ribosomal translation. Certainly, aminoglycosides interact with the decoding center of the ribosomal 16S subunit and cause misincorporation of an amino acid at the PTC site which allows translation to continue (Keeling and Bedwell, 2005; Zingman et al., 2007). RTC124 is believed to act at a different location on the ribosome (Linde and Kerem, 2008). It will be interesting to learn whether RTC#13 and #14 interact with the ribosome.

It has been reported that nonsense-mediated messenger RNA (mRNA) decay (NMD) can significantly affect RTC-induced PTC-readthrough because mRNA transcripts carrying nonsense mutations are degraded by this pathway (Wilkinson and Shyu, 2002; Holbrook et al., 2004). Therefore, inhibition of NMD may stabilize mutant mRNA transcripts and increase RTC-induced readthrough output (Linde et al., 2007; Linde and Kerem, 2008). We have examined this in our laboratory using many different ATM nonsense mutations, and we have failed to detect a clear pattern of NMD or inhibition of NMD, suggesting that NMD in A-T cells may be a matter of degree. NMD efficiency may also vary between different mutations and different genes. The role of NMD in RTC-induced treatment remains to be clarified.

**MATERIALS AND METHODS**

**Plasmids construction.** To construct plasmids containing stop codon mutations in their own context, we selected ATM nonsense mutations that resulted directly from disease-causing point mutations in A-T patients. The LCLS used in this study carried the following mutations: TAT51, homozygous 8977C→T (TGA A). Reverse-transcription PCR was performed using custom designed primers which introduced N- and C-terminal epitopes (c-myc and V5, respectively) into the PCR products (Du et al., 2008). The PCR products were then cloned into pDNAS/FRT/TORFTOPO plasmids according to the manufacturer’s protocols (Invitrogen). The PCR products for each mutant were mutagenized back to normal and used as paired normal control plasmids. DNA sequencing confirmed the PTT fragments in all constructs.

**High-throughput PTT-ELISA.** Three chemical libraries, ChemBridge, Prestwick, and MicroSource, were used in our primary screening. The final concentration for each compound tested was 10 µM. Screening was performed on a fully integrated CORE System (Beckman Coulter). PTT was performed in a reaction volume of 5 µl. The TNT T7 PCR Quick Master Mix, containing 20 µM methionine, was aliquoted into a 384-well low-volume plate (MatriCal Biosience). Compounds were added using a 384-well pin tool (V&P Scientific, Inc.). The PTT reaction was started by the addition of purified plasmid at 50 ng/well, which was incubated for 2 h at 30°C. PTT samples were subsequently stored at 4°C for ELISA analysis. For each plate, G418 at a final concentration of 1 µM served as positive control and reactions with 1% DMSO served as negative control. ELISA was performed in a 384-well MaxiSorp ELISA plate (Nunc). The plate was coated with 20 µl of 5 µg/ml mouse anti-Myc antibody (Invitrogen) overnight at 4°C, followed by washing with PBS and blocking with 50 µl PBSTM (PBS-containing 0.05% Tween-20 and 5% milk) for 30 min at 37°C. 20 µl H2O was added in PTT plate and 15 µl of reaction solution was transferred into ELISA plate, followed by overnight incubation at room temperature. After washing the plate, 20 µl of 1:500 mouse anti-V5 HRP (Invitrogen) was added and incubated for 2 h at 37°C. The plate was washed, incubated with 30 µl SuperSignal ELISA Pico working solution (Thermo Fisher Scientific), and measured on the Victor-3V using “top read” and an integration time of 0.5 s.

**Immunooassay for measurement of intranuclear ATM protein.** Readthrough-induced full-length ATM protein in A-T cells was measured by ATM immunooassay (Butch et al., 2004). Cell nuclear extracts were prepared using NE-PER protocol (Thermo Fisher Scientific). Nuclear extracts were analyzed by Western blot and AT-1981 IF. Immunostaining of nuclear foci of ATM-Ser1981 was performed as previously reported (Du et al., 2007). In brief, after being treated with compound for 4 d, cells were irradiated with 2 Gy and incubated at 37°C for 30 min. The cells were dropped onto coverslips, fixed with 4% paraformaldehyde, and permeabilized. Coverslips were blocked for 1 h and incubated with mouse anti-ATM pSer1981 for 1 h (1:500; Rockland Immunochemicals, Inc.). After a second blocking, cells were stained with FITC-conjugated anti–mouse IgG (1:150; Jackson ImmunoResearch Laboratories) for 1 h and mounted onto slides.

**RTC#13 and #14 induced a significant amount of dystrophin protein.** Results were consistent in three independent experiments.

**Figure 8.** RTCs restored full-length dystrophin protein in mouse *mdx* myotubes (TAA). Cells were prepared and treated with RTCs for 3 d. Dystrophin proteins were detected by Western blot analysis using immunoprecipitation. Wild-type cells were used in the experiment to localize dystrophin protein. Gentamicin was used as a positive readthrough control. Both RTC#13 and #14 induced a significant amount of dystrophin protein. Results were consistent in three independent experiments.
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