Ultrashort and progressive 4sU-tagging reveals key characteristics of RNA processing at nucleotide resolution

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RNA synthesis and decay rates determine the steady-state levels of cellular RNAs. Metabolic tagging of newly transcribed RNA by 4-thiouridine (4sU) can reveal the relative contributions of RNA synthesis and decay rates. The kinetics of RNA processing, however, had so far remained unresolved. Here, we show that ultrashort 4sU-tagging not only provides snapshot pictures of eukaryotic gene expression but, when combined with progressive 4sU-tagging and RNA-seq, reveals global RNA processing kinetics at nucleotide resolution. Using this method, we identified classes of rapidly and slowly spliced/degraded introns. Interestingly, each class of splicing kinetics was characterized by a distinct association with intron length, gene length, and splice site strength. For a large group of introns, we also observed long lasting retention in the primary transcript, but efficient secondary splicing or degradation at later time points. Finally, we show that processing of most, but not all small nuclear (sn)RNA-containing introns is remarkably inefficient with the majority of spliced/degraded introns. In summary, our study yields unparalleled insights into the kinetics of RNA processing and provides the tools to study molecular mechanisms of RNA processing and their contribution to the regulation of gene expression.

[Supplemental material is available for this article.]
high-quality sequencing data. The combination of ultrashort and progressive 4sU-tagging from 5- to 60-min labeling time then allows unparalleled insights into the kinetics of RNA processing, in particular RNA splicing and processing of noncoding RNAs.

Results

Ultrashort 4sU-tagging is compatible with RNA-seq in human B-cells

Newly transcribed RNA obtained by 4sU-tagging contains substantially greater amounts of large, unprocessed transcripts than regularly found in total cellular RNA. This is readily visualized by electrophoretic analysis (Do¨lken et al. 2008). When shortening the duration of 4sU-tagging the average age of nascent transcripts in newly transcribed RNA decreases. We thus hypothesized that RNA-seq combined with progressive reduction of the duration of 4sU-tagging could be employed to study the kinetics of RNA processing. For this purpose, we performed a time course experiment of 4sU-tagging in DG75 human B-cells consisting of five samples with 60, 20, 15, 10, and 5 min of 4sU-tagging. At the end of 4sU exposure, cells were harvested using TRIzol, total cellular RNA was prepared, and newly transcribed RNA was purified. The relative abundance of newly transcribed, tagged RNA in total cellular RNA decreased from 3.5% of total RNA after 1-h 4sU-tagging to ~0.8% after 5 min (Fig. 1A). Newly transcribed RNA from all five labeling conditions was subjected to RNA-seq analysis using sequencing by ligation (SOLiD II, Applied Biosystems). In the following, we will refer to these samples as ‘5-min 4sU-RNA’ to ‘60-min 4sU-RNA.’ In addition, total and untagged RNA following 60 min of 4sU-tagging were sequenced. As newly transcribed RNA may not yet be poly-adenylated, no poly(A) selection was performed. For each sample, between 2.8 and 4.4 million reads could be uniquely mapped by aligning them first to the human tran-
Progressive 4sU-tagging reveals RNA processing

Intron decay is correlated to intron length, gene length, and splice site strength

Interestingly, the decrease in the number of exon–intron junction reads appeared to be delayed compared with the decrease in the intronic reads themselves (Fig. 1C). As the relative contribution of long introns to the number of intronic reads is higher than to the number of exon–intron reads (Supplemental Fig. 1A), we hypothesized this to be due to a faster decay of long introns. In this case, intronic read numbers, which are disproportionately determined by long introns, would decrease faster with increasing labeling time than exon–intron junction read numbers. Indeed, no delay could be observed when read numbers were analyzed selectively for three approximately equally sized groups of introns with similar length (Supplemental Fig. 1B–D).

To quantify the rate of decay for each intron, ratios of intron reads in 60-min 4sU-RNA compared with 5-min 4sU-RNA (60/5 min ratios) were calculated and correlated with intron length. Thus, high 60/5 min ratios correspond to low decay rates and vice versa. On the individual intron level, a correlation with intron length was basically nonexistent (Spearman rank correlation \( r_s = -0.037, P\text{-value} = 0.0039 \)). However, when introns were binned according to length and average 60/5 min ratios were calculated for each bin, a very clear trend was observed (Fig. 1D). Interestingly, the highest 60/5 min ratios, i.e., the lowest decay rates, were not observed for the shortest introns, but for introns in a range of \( 300–400 \) nucleotides (nt). Decay rates increased both for introns longer as well as shorter than this range, with the highest decay rates observed for introns shorter than \(-1500 \) nt. Accordingly, after normalizing intronic read counts to intron length, the lag between the loss of intronic reads and exon–intron junction reads over time disappeared (Fig. 1E).

To investigate whether these findings were influenced by any bias introduced by our experimental setup or computational analysis, we correlated 60/5 min ratios and intron length with other intron features. While there was no significant correlation between the relative position of an intron in a gene and its 60/5 min ratio or length, gene length was correlated to both characteristics (\( r_s = -0.24 \) for 60/5 min ratios and \( r_s = 0.38 \) for intron length, \( P\text{-value} < 10^{-15} \) in both cases). Here, the correlation between gene length and 60/5 min ratio was very clear when binning according to gene length (\( r_s = -0.94 \)): As average gene length increased, average 60/5 min ratios decreased. An analysis of 60/5 min ratios binned first according to gene length into three groups, and then for each group binned according to intron length, indicated that the contributions of gene and intron length were mostly independent of each other (Supplemental Fig. 1E). Irrespective of gene length, intron length and 60/5 min ratios showed the same characteristic correlation with a peak of 60/5 min ratios in the low-to-medium intron length range (300–400 nt) and decreasing ratios on either side.

Finally, we investigated whether splice site strength had any effect on splicing kinetics. Splice site strength was quantified in terms of splice site scores calculated with MaxEntScan (Yeoe and Burge 2004). Similar to intron length, there was only an extremely weak correlation between splice site scores (\( 5^\prime \) only, \( 3^\prime \) only, and \( 5^\prime + 3^\prime \) scores) and 60/5 min ratios (\( r_s = -0.055, -0.044, \) and \(-0.072, \) respectively) or intron length (\( r_s = 0.055, 0.024, \) and \(0.046, \) respectively). However, when binning introns according to the splice site strength, there was a trend toward higher 60/5 min ratios (slower splicing kinetics) for very weak splice sites (Supplemental Fig. 1F). This effect was mostly independent of intron length (Supplemental Fig. 1G).

Distinct classes of introns are defined by their splicing kinetics

To further investigate differences in the kinetics of intron processing, we first focused on the most highly expressed genes as intron expression levels in newly transcribed RNA, although substantially higher than in total RNA, are much lower than the expression levels of the surrounding exons. This is due to the large fraction of introns (\( >65\% \)) already spliced and decayed in 5-min 4sU-RNA. Expression levels of genes were quantified in terms of reads per kilobase of gene per million mapped reads (RPKM) after normalizing for mappability (see Methods), and the analysis was focused on genes with an RPKM \( \geq \) 11 in all RNA samples (525 genes). Since ribosomal proteins are generally highly expressed, they, as well as other genes involved in translation and ribosome biogenesis, were enriched in this set. Other enriched functions included RNA splicing, ATPase activity, cell cycle, and regulation of apoptosis (see Supplemental Table 1 for a full list). For these genes, we distinguished between introns absent (RPKM < 0.5: 1014 introns, Supplemental Table 2) or present (5838 introns, Supplemental Table 3) in 5-min 4sU-RNA. Even after excluding 50 absent introns (\( \approx 5\% \)) that were shorter than the read length (35 nt) and, thus, could not contain any intronic reads, absent introns were significantly shorter than the present ones (Wilcoxon test, \( P\text{-value} < 10^{-15} \)). Furthermore, they were located closer to the 3′ end of the gene than present introns (Wilcoxon test, \( P\text{-value} = 0.0042 \)) with 12% of the absent introns being the last intron of the gene compared with 7% for present introns (Fisher's exact test, \( P\text{-value} < 10^{-6} \)).
This suggests that at least some of these introns were part of longer transcript versions that were not transcribed in this form in the DG75 cells. For other introns, possible explanations for their absence in 5-min 4S-U-RNA might be (1) very fast co-transcriptional splicing, (2) problems in sequencing, or (3) problems in mapping these parts of the pre-mRNA, e.g., due to repetitive sequences. Interestingly, in many absent introns, both neighboring exons were well expressed and precisely delimited. This indicates rapid co-transcriptional splicing and intron degradation rather than sequencing bias. In addition, there was no significant increase for the absent introns in the frequency of repetitive sequences identified by RepeatMasker (Smit et al. 1996) or the frequency of nonunique read mappings. Notably, the fraction of absent intron positions contained within repetitive sequences was actually significantly smaller than for present ones (Wilcoxon test, \( P \)-value < 10\(^{-4}\)). These analyses confirm that numerous transcripts in 5-min 4S-U-RNA (>65%) had already been spliced and their introns had been degraded.

Remarkably, exon–intron junction reads for these absent introns were regularly observed, although only at ~60% of the level of introns present in 5-min 4S-U-RNA (Wilcoxon test, \( P \)-value < 10\(^{-18}\)). The frequency of exon–exon junction reads was reduced to a similar degree. This indicates that the proportion of completed splicing events including the fusion of neighboring exons was not significantly higher than for present introns. A possible explanation for this observation is that major parts of the absent introns are already degraded while the exon–intron junctions are still connected to the splicing machinery in a state prior to exon fusion.

An example for such an absent intron is shown for the RPL23A gene (Fig. 2A). In this case, the absent intron is surrounded by a retained intron contained in an alternative transcript of RPL23A. According to the Ensembl annotation, this alternative transcript is not translated. It is important to note that the absence of this intron is not due to repetitive sequences in this region, which might result in nonuniquely mapped reads and their subsequent exclusion during mapping. Although two repeats were identified by RepeatMasker in this absent intron, sequence divergence to the corresponding repeat consensus was rather high (6.2% and 11.3%, respectively). Accordingly, only one additional nonuniquely mapping read was found in all samples. This confirms that the low expression of this intron is not an artifact created by the removal of nonunique mappings. Interestingly, the intron retention in RPL23A is still observed at considerable levels in 60-min 4S-U-RNA, but hardly detectable in total and untagged RNA. This suggests that this alternative splicing variant is erroneously produced at considerable levels but quality control mechanisms of the cell later on result in its delayed but nevertheless highly efficient removal. This may either result from decay of the whole transcript or secondary splicing events removing only the retained intron.

To systematically mine for such and other distinctive kinetics of intron processing, we used cluster analysis on the introns still present in 5-min 4S-U-RNA. Introns were clustered according to their relative abundance compared with the expression level of the gene (intron/gene ratio). The intron/gene ratio approximately represents the fraction of pre-mRNAs or transcripts still containing the intron. A stable clustering was obtained by repeated k-means clustering (see Methods). The resulting 20 largest clusters represented 85% of introns (Supplemental Fig. 2A). Although distinct differences between introns of the same gene were observed, they were significantly enriched in the same clusters (Fisher's exact test, \( P \)-value < 10\(^{-13}\)). This indicates that introns of the same genes tend to have similar splicing kinetics.

A visual inspection of the clusters identified distinctive subgroups of clusters with different absolute 5-min intron/gene ratios but similar trends across the time course, i.e., similar relative ratio changes over time. For random data sets analyzed as controls, no distinctive subgroups were observed (Supplemental Fig. 2B). To systematically identify such subgroups of splicing patterns, the 20 largest clusters were clustered again after normalizing the intron/gene ratios to 5-min 4S-U-RNA. Thus, exon/intron ratios in 5-min 4S-U-RNA were set to 100%. This resulted in four distinct classes of intron splicing (Fig. 2B; Supplemental Table 3). The largest class, Class 1, representing 3908 introns, was characterized by an almost linear slope in intron/gene ratios up to 20-min 4S-U-RNA. Class 2 (580 introns) showed a smaller slope than Class 1, suggesting a lower splicing rate. In contrast, Class 3 (338 introns) was characterized by a more rapid decrease in intron/gene ratios between 5- and 10-min 4S-U-RNA, indicating a faster splicing rate. Finally, Class 4 (109 introns) showed a very interesting pattern. Here, intron/gene ratios remained remarkably stable from 5- to 60-min 4S-U-RNA, but then dropped significantly some time later to reach similar low levels in total and untagged RNA as observed in the other classes. Exemplary introns for Class 1 and 4 are shown in Figure 2C,D. Please note that the division into different classes was not always clear-cut. For instance, although cluster 3 (238 introns) is assigned to Class 2, it shows a similar trend as Class 4 introns with stable intron levels up to 20–60 min and subsequent decay. In general, Class 2 constitutes a weaker version of the delayed decay in Class 4, while Class 3 contains extreme cases of rapid intron removal and decay.

**Monitoring the fate of alternative splicing products**

Interestingly, Class 4 introns and to a lesser degree Class 2 introns showed a pattern very similar to the known intron retention example shown in Figure 2A. This suggests that these introns likely constitute novel alternative splicing events resulting in retained introns. To provide evidence for this hypothesis, degradation patterns for known alternative splicing products were investigated. For this purpose, exon boundaries were recalculated based on Ensembl transcripts and the redefined exons were classified either as core exons, retained introns, or alternative 3’ or 5’ exon ends (see Supplemental Material). Exon expression levels were divided by the overall gene expression level (exon/gene ratios) and normalized to 5-min 4S-U-RNA (Fig. 2E). As expected, core exon/gene ratios were stable across all time points. In contrast, for retained introns the same pattern was observed as for Class 4 and some Class 2 introns. The same pattern was also seen for alternative 3’ and 5’ ends, although much less pronounced. This implies that alternative transcripts containing retained introns are present at substantial numbers (>30%) until 60-min 4S-U-RNA but are subsequently degraded quite rapidly, e.g., by nonsense-mediated decay or by secondary splicing events.

**Properties of splicing classes**

Intron length, gene length, and splice site strength were analyzed to reveal their contribution in defining the kinetics of splicing in the four classes of introns. Interestingly, significant differences in the distribution of these features were observed (Fig. 3A). Class 1 introns were on average longer (by 36%, Wilcoxon test, FDR corrected \( P \)-value < 10\(^{-12}\)), contained in longer genes (by 76%, \( P \)-value < 10\(^{-15}\)), and showed increased splice site strength (by 4.5%, \( P \)-value < 10\(^{-13}\)) compared with introns of the other classes. As all of these
Figure 2. Characterization of distinct intron splicing kinetics. (A) Decay of an alternative splicing isoform of the RPL23A gene. Read densities for 5- to 60-min 4sU-RNA as well as total and untagged RNA are shown in shades of gray. This figure and all others showing read densities were created using the UCSC Genome Browser (Kent et al. 2002). The corresponding UCSC Genome Browser session containing read density values for all 525 genes analyzed can be accessed via http://www.bio.ifi.lmu.de/en/4sU-seq. Exons are indicated by boxes, introns by lines. Regions that are part of noncoding transcripts are indicated by boxes of smaller height. The known retained intron not translated is additionally indicated in the last line in black. The intron marked in gray is absent in all 4sU-RNA samples (RPKM < 0.5), which is indicative of very fast splicing. In total and untagged RNA, the retained intron is also absent, indicating either nonsense-mediated decay of the alternative splicing product or a secondary splicing event. (B) Higher level clustering of the intron clusters identified in the first clustering step (see also Supplemental Fig. 2). Clustering was performed based on intron/gene ratios of cluster representatives normalized to 5-min 4sU-RNA levels. Four classes of intron processing were identified: (Class 1) “Normal” decay with a linear slope in the first 20 min; (Class 2) reduced decay rates; (Class 3) increased decay rates; (Class 4) retained introns that are stable during the first 60 min but are eventually spliced and/or degraded. (C) Read density in all samples for an exemplary Class 1 intron. (D) Read density for an exemplary Class 4 intron. (E) Exon/gene ratios in all samples for core exons, exons constituting retained introns, as well as alternative 3’ and 5’ ends. Again, exon/gene ratios were normalized to ratios in 5-min 4sU-RNA.
factors were positively correlated with intron decay rates, it is not surprising that these introns are lost relatively fast compared with the majority of other introns (Classes 2 and 4). Strikingly, Class 3, which was characterized by the most rapid intron decay, contained shorter introns and shorter genes than Class 1, while splice site strength was neither significantly increased nor decreased. In contrast, Classes 2 and 4, which also contained significantly shorter introns, were characterized by both significantly shorter gene length and weaker splice sites compared with Classes 1 and 3 (Wilcoxon test, \( P \)-value < 10^{-5}). These results suggest that for short introns, gene length and splice site strength make a marked difference, resulting in either very fast or very slow splicing and intron decay. In particular, reduced splice site strength may result in the retention of introns (resulting in Class 4 or Class 2 introns) due to poor recognition of splice sites by the splicing machinery. For the latter two classes, the strongest difference was in intron length with Class 2 having significantly longer introns than Class 4. Thus, the four classes are characterized by distinct combinations of intron length, gene length, and splice site strength.

Characteristics of snoRNA processing

Interestingly, both Class 2 and Class 4 were significantly enriched for small nuclear RNA (snoRNA) precursor introns. SnoRNAs are small noncoding RNAs mainly involved in the chemical modification of other noncoding RNAs, in particular rRNAs. They are mostly encoded within introns of protein-coding or noncoding genes and are excised from these larger RNAs during splicing (Hirose et al. 2003; Dieci et al. 2009). In our case, 22 out of 580 Class 2 introns (3.8%, Fisher’s exact test, FDR corrected \( P \)-value = 0.0027) and seven out of 109 Class 4 introns (6.4%, Fisher’s exact test, FDR corrected \( P \)-value = 0.011) contained snoRNAs. In contrast, snoRNA-containing introns were underrepresented in Class 1 (52 out of 3908 introns, 1.3%, \( P \)-value = 0.00024). To analyze this observation in more detail, the RNA-seq data were used to investigate processing of snoRNA transcription units. Almost all human snoRNAs (>90%) are processed from introns with only a single snoRNA generated from each intron (Dieci et al. 2009). For these snoRNAs, splicing is generally required to make the 5' and 3' ends of the snoRNA-containing intron accessible to trimming by nuclear RNAses. As the majority of snoRNA-containing genes were found to be among the most highly transcribed genes, this group of rather small noncoding RNAs was ideally suited for further analysis. Again, we focused on the 5838 well expressed introns of which 121 contained snoRNAs. Of these snoRNA precursor introns, 88 (72%) were assigned to one of the four classes. The remaining ones were not assigned to the 20 largest clusters. Please note that in this and all previous analyses reads mapping to snoRNAs were assigned only to the snoRNAs and not used for calculating the intron levels.

As Classes 2 and 4 showed reduced intron decay rates, we wondered whether this might be a general feature of snoRNA-containing introns. Indeed, intron/gene ratios of snoRNA precursor introns were significantly increased compared with all other introns at all times. (C) Read density plot illustrating the processing of an exemplary precursor intron to the mature snoRNA. One can observe both the substantially slower decay of the intronic sequences surrounding the mature snoRNAs, which was typical for intronic snoRNAs, and the characteristic 5' read start pattern of mature snoRNAs in total and untagged RNA.
Accordingly, snoRNA intron processing generally appears to be a continuous process and not a case of temporary intron retention and delayed processing/decay. Remarkably, splice site scores of snoRNA-containing introns were not significantly reduced compared with other introns. This suggests that the reduced decay rates are not due to poor recognition of the splice site by splicing factors. In contrast, both intron length and length of the corresponding genes were reduced on average by ~40% (Wilcoxon test, P-value = 0.0005) and 60% (P-value < 10^{-4}), respectively. As both short-to-medium intron length and short gene length have been found to be associated with reduced intron decay rates, even if snoRNA encoding introns were excluded from the analysis, this may provide a possible explanation for the smaller splicing rates of snoRNA introns.

One representative example of a snoRNA precursor intron is shown in Figure 3C. Here, the SNORD4B snoRNA is excised from an intron of the RPL23A gene. In this case, large parts of the precursor intron were clearly present until 60-min 4sU-RNA, whereas only the mature snoRNA was detected in total and untagged RNA. Interestingly, the mature snoRNAs can be identified by reads starting predominantly at the 5' end of the snoRNA. These sequences most likely represent cloning products of mature snoRNAs not affected by the fragmentation procedure applied during library preparation. Any fragmentation of these small noncoding RNAs probably resulted in the loss of the resulting snoRNA fragments as the standard SOLiD protocol includes a size selection during library preparation to remove cDNAs without inserts.

Due to this apparent cloning/sequencing bias, determination of snoRNA stability using the RNA-seq data is not possible. Thus, we analyzed snoRNA half-lives obtained using Affymetrix Gene ST 1.0 microarrays in DG75-eGFP and two other human B-cell lines (DG75-10/12 and BCBL-1) (Dölen et al. 2010) based on newly transcribed/total RNA ratios. Surprisingly, snoRNAs were highly enriched among the most short-lived transcripts in all three B-cell lines in the microarray data (Fig. 4A). Here, snoRNAs represented >20 of the 40 most short-lived transcripts in all three human B-cell lines. This was surprising as these small noncoding RNAs
are involved in metabolic processes, which are usually associated with long-lived transcripts (Friedel et al. 2009). In addition, there is so far no evidence for snoRNAs being used up and metabolized while exerting their function in rRNA maturation. Furthermore, snoRNAs are small RNA molecules (60–300 nt; Kiss 2002). Therefore, they are prone to reduced capture rates during 4sU-tagging due to their very low uridine (and thus low 4sU) content (Friedel et al. 2009; Miller et al. 2009). Capture rates can be assessed by comparing the log2 (newly transcribed/total RNA) ratios with the uridine content of transcripts (Fig. 4B). As expected, average log ratios decreased for very short transcripts with very low uridine content. For virtually all snoRNAs, however, these ratios were not only not reduced, but rather surprisingly high for transcripts of such short length. If we had corrected for size-dependent reduced capture efficiency using computational methods (Miller et al. 2009, 2011), these ratios would have been even higher, resulting in even shorter RNA half-lives.

Remarkably, the apparent short snoRNA half-lives from microarrays could also not be verified by Northern blot analysis. Here, snoRNA levels for eight short-lived, seven medium-lived, and three long-lived snoRNAs (according to the microarray data) were determined following 6-h transcriptional arrest with either Flavipiridol (FL) or Actinomycin D (Act-D). The very short-lived c-myc transcript and the long-lived glyceraldehyde-3-phosphate dehydrogenase (GAPDH) served as controls. While a reduction in snoRNA levels following transcriptional arrest was observed for three snoRNAs (SNORD22, SNORA38B, and SNORA73A), expression of the majority of snoRNAs remained unaltered (Fig. 4C). Although a rapid stabilization of these snoRNAs due to transcriptional arrest cannot be ruled out completely, the alterations were much smaller than for other short-lived RNAs (e.g., MYC or FOS) known to be rapidly stabilized after transcriptional arrest. We conclude that most mature snoRNAs molecules are not highly unstable, in contrast to what was implied by the microarray data. However, as the majority of snoRNAs are well expressed in human B-cells and were highly consistent in three different cell lines, these measurements are very unlikely to be due to technical problems or artifacts of the microarray measurements.

As microarray probes cannot distinguish between mature snoRNAs and their precursors, we hypothesized that microarrays measured the short RNA half-lives of instable snoRNA precursors, rather than of the mature snoRNAs. Accordingly, the very short half-lives observed for many “snoRNA” probe sets in the microarray data likely represent the half-lives of their parental introns. Thus, they correspond to rapid splicing and decay of the parental introns, which are degraded rather than processed to a (stable) mature snoRNA. In conclusion, the short snoRNA half-lives observed by microarray analysis actually reflect inefficient processing of many snoRNAs, i.e., only a small fraction of introns is actually processed to mature snoRNAs while the majority of introns are simply spliced and degraded. Interestingly, very short microarray-derived half-lives were observed only for snoRNAs encoded within introns of noncoding and protein-coding genes, but not for snoRNAs with independent promoters, which provides further support to this conclusion (Fig. 4D).

In contrast to most other snoRNAs, snoRNAs of the SNORD116 cluster were found to possess rather long RNA half-lives (>5 h) in the microarray data (Fig. 4D). This would be consistent with substantially more efficient snoRNA processing. It is important to note that their long RNA half-lives were not due to low signal intensities of the corresponding probe sets. These snoRNAs all derive from introns of the SNURF–SNRPN transcription unit (Runte et al. 2001), which is expressed from the imprinted SNURF–SNRPN domain. It encodes for at least two long, paternally expressed noncoding RNAs, which both encode for multiple intronic snoRNAs (Royo and Cavaille 2008; Vitali et al. 2010). Little is known about their biogenesis and function. The second cluster of snoRNAs (SNORD115) encoded in this domain is brainspecific (Cavaille et al. 2000) and was hardly expressed at all in the B-cells under study. Although found only at relatively low levels in our RNA-seq data, the mature snoRNAs of the SNORD116 cluster were expressed at greater levels than the surrounding intronic sequences (Fig. 5A). Furthermore, reads derived from the SNORD116 cluster generally started at the 5′ end of the snoRNA even in 5-min 4sU-RNA (Fig. 5B), which is in stark contrast to the other intronic snoRNAs (Fig. 5C). This strongly indicates that these reads derive from mature snoRNAs. Accordingly, most SNORD116 snoRNAs were already fully processed in 5-min 4sU-RNA, which indicates highly efficient snoRNA processing within the SNORD116 cluster.

Figure 5. Processing of the SNORD116 cluster. (A) Read density for the SNORD116 cluster region (overview). (B) Read density plot for a typical snoRNA of the SNORD116 cluster only shows reads at the 5′ end of the mature snoRNA, but no intronic sequences, thus representing sequencing of the mature, unfragmented snoRNA. (C) Read density plot for a typical intronic snoRNA. Here, intronic sequences were observed until 60-min 4sU-RNA.
Discussion

Metabolic labeling of newly transcribed RNA with 4-thiouridine (4sU-tagging) is superior to simply analyzing total cellular RNA and allows distinguishing alterations in RNA synthesis and decay (Döldken et al. 2008; Friedel et al. 2009; Rabani et al. 2011). In this study, we show that characteristics of RNA processing can be examined by performing ultrashort and progressive 4sU-tagging combined with RNA-seq. In human B-cells, as little as 5 min of 4sU-tagging provided sufficient amounts of newly transcribed RNA to perform RNA-seq. Previously, 4sU-tagging was shortened to 10–15 min but only combined with microarray (Döldken et al. 2008) and nCounter technology (Rabani et al. 2011) as not enough RNA material was thought to be recoverable for sequencing. RNA-seq was only performed following much longer labeling (e.g., 2 h) (Schwanhäusser et al. 2011). In order to shorten the duration of 4sU labeling down to 5 min and still obtain sufficient newly transcribed RNA for next-generation sequencing, we increased the amount of total RNA with which we started from 70 to 150 μg. In addition, we benefited from a slightly increased efficiency of 4sU uptake by cells growing in suspension (B-cells) compared with adherent cells (murine fibroblasts). As 4sU incorporation is strongly dependent on the employed 4sU concentration, this approach is readily adaptable to other cell types by simply increasing the 4sU concentration for ultrashort labeling.

The high purity of the newly transcribed RNA samples was confirmed by the much higher percentage of intronic reads and exon–intron junction reads inversely correlated with the duration of labeling. Interestingly, our data showed that >65% of intronic sequences had already been spliced and decayed in 5-min 4sU-RNA, which highlights the need to perform ultrashort 4sU-tagging. To further characterize processing of introns, we analyzed the relative loss of intronic reads over time (5-min 4sU-RNA to 60-min 4sU-RNA) to define splicing kinetics for >$800 introns and correlated this with various features. Remarkably, intron decay rates were found to be influenced by both intron and gene length. The correlation to intron length was mostly independent of gene length as it was also observed for genes of similar length. As these observations were based on the comparison of read counts in 60-min 4sU-RNA to 5-min 4sU-RNA, it is unlikely that they are due to sequencing artifacts or insufficient length normalization. These problems would affect intron read counts in both samples to a similar degree and, thus, they would not be relevant when calculating ratios of read counts for each intron. In conclusion, although we cannot completely exclude a minor bias introduced during sample and library preparation negatively affecting sequence reads derived from very large genes/pre-miRNAs, our data indicate that any bias introduced did not substantially affect the major findings of this study.

Interestingly, numerous introns were already absent in 5-min 4sU-RNA. Although some of these probably represented non-transcribed variants at the 3′ end of a gene and some may simply reflect sequencing bias, a considerable number were surrounded by well-expressed exons, suggestive of very fast splicing. Remarkably, we still observed exon–intron junction reads for these absent introns albeit at a lower frequency than for introns present in 5-min 4sU-RNA. Furthermore, the number of exon–exon junction reads was not increased compared with the number of exon–intron junction reads. This hints at degradation of these introns concurrently with splicing, which would be consistent with previous results showing that tethering of exons to the RNA polymerase II results in correct splicing of these exons even though the intron connecting them may no longer be continuous, e.g., due to ribozyme-mediated cleavage and partial degradation (Dye et al. 2006).

When looking more closely at the kinetics of splicing, we identified clusters of introns spliced with distinct kinetics accompanied by an enrichment for similar splicing kinetics within a given gene. It is important to note that the reported splicing kinetics were also observed for less abundant genes with more diverse functions. When classes were extended by so far unclassified present introns (RPKM ≥ 0.5) of genes with a lower minimum expression value (RPKM ≥ 5) (2468 genes; 29,947 introns), the same trends were observed for each class as before (Supplemental Fig. 2C). The most interesting kinetics were observed in introns of Class 4 and less pronounced in Class 2. These showed relative high intron levels even in 60-min 4sU-RNA—indicative of retained introns. This was associated with reduced splice site strength most likely resulting in missed splice site recognition contributing to intron retention. Nevertheless, intron levels dropped substantially in total and untagged RNA. This is consistent with a delayed but nevertheless efficient removal of these intronic sequences. Many known and at least 100 novel alternatively spliced transcripts containing retained introns showed this pattern. This temporally delayed but nevertheless eventually efficient removal of retained introns either may be due to RNA degradation by nonsense-mediated RNA decay or may represent novel secondary, i.e., post-transcriptional splicing events. In principle, the two could be distinguished, as secondary splicing events would only remove the initially retained introns while nonsense-mediated decay would also affect levels of the surrounding exons and thus overall transcript RNA half-lives. Unfortunately, the fraction of transcripts of a gene still containing retained introns in 5-min 4sU-RNA was not much larger than 30% on average. Thus, the effect on half-lives was not large enough to differentiate the two alternatives based on the current data. Further studies will thus need to employ knockout/down approaches targeting nonsense-mediated RNA decay to answer this interesting question. Nevertheless, both alternatives suggest highly effective cellular quality control measures to ensure correct splicing, as the level of transcripts containing retained introns in total and untagged RNA was very low.

Another interesting finding of this study was the poor processing efficiency of many but not all human snoRNAs. In microarray data of newly transcribed/total RNA ratios derived from 1-h 4sU-tagging (Döldken et al. 2010) snoRNAs seemed to be the most short-lived cellular transcripts in three human B-cell lines. Northern blot analysis of mature snoRNAs after 6 h of transcriptional arrest clearly excluded such short RNA half-lives for most of the mature snoRNAs, thereby at first contradicting the microarray-based measurements. However, as microarray probe sets cannot differentiate between a small mature snoRNA and its much larger precursor, these “seemingly” short RNA half-lives only reflect inefficient processing of snoRNAs from their much larger precursors, i.e., degradation of the parental introns without processing to the mature snoRNAs. Poor processing efficiency of snoRNAs derived from intronic sequences indicates competition between splicing factors (resulting in subsequent intron degradation) and the snoRNA processing machinery. Evidence for this hypothesis is provided by the observation that introns containing snoRNAs are spliced and degraded much slower than other introns. It is tempting to speculate that the low basal processing efficiency of many snoRNAs may offer the opportunity for significant regulation of snoRNA expression levels by modifying their processing efficiency. Indeed, up-regulation of snoRNA levels has recently been reported for cells expressing a mutant form of the carboxy-terminal
domain (CTD) of the large subunit of RNA polymerase II (Sims et al. 2011). Expression of a CTD mutant deficient in arginine methylation resulted in a significant increase of steady state levels of a variety of snoRNAs and snRNAs (small nuclear RNAs), while the levels of all other categories of RNAs, e.g., mRNAs, remained unaffected. It will be interesting to test whether this mutant shows alterations in RNA splicing, which might account for the alterations in snoRNA levels. An alternative explanation for the large number of snoRNA-containing introns being spliced and degraded rather than processed into mature snoRNAs would be that the expression levels of snoRNA-binding proteins may not be sufficiently high to bind all newly synthesized snoRNA precursors and thereby prevent their degradation. In this case, snoRNA levels would not be defined transcriptionally but by the abundance of their respective snoRNA-binding proteins.

Interestingly, RNA processing of intronic snoRNAs was not always found to be inefficient as exemplified by the majority of snoRNAs of the SNORD116 cluster. Although we did observe weak expression of the intronic regions surrounding the snoRNAs, accumulation of reads derived from the 5′ end of the mature snoRNAs in all samples provided evidence that mature snoRNAs were already generated within the first few minutes of synthesis and thus became apparent even in 5-min 4sU-RNA. This also indicates that snoRNA processing can indeed be both fast and efficient. For other snoRNAs with less efficient processing, this clear bias of read starts characteristic for mature snoRNAs was only observed later on, indicating that the majority of their parental introns are not rapidly spliced and processed to snoRNAs. The underlying molecular mechanism of this substantial difference in processing efficiency of the paternally imprinted SNORD116 snoRNAs remains to be elucidated.

RNA processing involves numerous complex and highly regulated molecular processes. This field of research has recently received a dramatic boost by the development of high-throughput technology for studying RNA–protein interactions at nucleotide resolution. Both HITS-CLIP and PAR-CLIP can now be employed to reveal thousands of protein–RNA interaction sites in a single experiment and thus serve to unravel the underlying functional networks and molecular mechanisms (Ule et al. 2003, 2006; Hafner et al. 2010). Our approach now provides the means to directly study the functional consequences of these RNA–protein interactions on RNA processing. In this study, we restricted our approach to RNA splicing and the processing of the highly abundant snoRNAs. With more in-depth sequencing and sequencing of shorter RNAs, RNA processing of other noncoding RNAs, such as miRNAs or lincRNAs, could also be studied in detail. One of the most important conclusions of this work is that RNA processing efficiency needs to be considered when studying the regulation of RNA expression levels. RNA processing is likely to provide a notable contribution to the regulation of both large and small noncoding RNAs. Application of RNA-seq combined with ultrashort and progressive 4sU-tagging will thus dramatically enhance our understanding of the underlying molecular mechanisms and regulatory networks.

**Methods**

**Cell culture and 4sU-tagging**

DG75-eGFP human B-cell lines were cultured in RPMI 1640 medium containing 10% fetal calf serum, 100 IU/mL Penicillin, 100 μg/mL Streptomycin, and 2 mM L-Glutamin. Newly transcribed RNA was labeled for 5, 10, 15, 20, or 60 min using 500 μM 4sU. Total cellular RNA was prepared from cells using TRIzol reagent (Invitrogen) following the modified protocol by Chomczynski and Mackey (1995). Newly transcribed RNA was purified from 150 μg (5- to 15-min 4sU), 100 μg (20-min 4sU), and 70 μg total RNA (60-min 4sU). Separation of total RNA into newly transcribed and untagged preexisting RNA was performed as described (Dölden et al. 2008) using 100 μL streptavidin beads (Miltenyi Biotec).

**Next-generation sequencing**

RNA sequencing was performed using the sequencing-by-ligation SOLiD II platform (Applied Biosystems). A modified whole-transcriptome analysis (WTAK) protocol was employed. Three-hundred nanograms of RNA was used in the library construction process. Volume of the samples was adjusted by vacuum centrifugation, as concentrations of the early newly transcribed RNA samples tended to be low after depletion. Integrity, size distribution, and yield were monitored after RNaseIII fragmentation (10 min, 37 °C) on an Agilent RNA 6000 PicoChip. If newly transcribed RNA samples resulted in larger than expected fragments, samples were redigested using RNaseIII for additional 2 min at 37 °C, repurified, and reordered as described above. Libraries of individual samples were prepared using barcoded adaptors and sequenced on the SOLiD instrument using standard protocols. No deviation in quality or length of the newly transcribed RNA sequences was observed when comparing with other standard RNAs.

**Read mapping**

Reads were mapped in a three-step process using the Bowtie alignment program (Langmead et al. 2009). First, reads were aligned to pre-rRNA sequences (18S, 5.8S, 28S, and spacer regions). The remaining unmapped reads were aligned to all Ensembl transcripts (Ensembl version 60) excluding pseudo-genes and haplotypes to identify exonic and exon–exon junction reads (aligned reads overlapping an exon–exon junction by ≥1 nt). Reads that remained unmapped after step two were aligned to the human reference genome (GRCh37/hg19) to identify intron and exon–intron junction reads (overlapping an exon–intron junction by ≥1 nt). For each read the best alignment location was used. Reads mapping equally well to two different locations were discarded. The following Bowtie settings were used: seed region = first 20 nt, three mismatches allowed in the seed, five in the whole alignment.

**Quantification of gene, exon, and intron expression levels**

Expression levels of genes, exons, and introns were estimated using the standard RPKM measure (number of reads per kilobase of gene, exon, or intron per million mapped reads) (Mortazavi et al. 2008). Number of reads mapping to a gene was determined as the total number of exons and exon–exon junction reads for this gene. To calculate RPKM values for exons and introns respectively, only reads mapping completely within this region were used. To account for problems in mapping reads to repetitive sequence regions, the effective length of exons and introns was used instead of the actual length. The effective length was calculated in the following way. First, in silico reads were simulated by sliding a window across gene regions with the size of the read length in the experiment (35 nt). Thus, the simulated read set contains exactly one read from each position in each gene. The simulated reads were then mapped using the same three-step procedure described above. The effective length was then defined as the number of positions within the respective region (exon, intron, or gene) which
had exactly one correctly and uniquely mapped read starting at this position.

Intron clustering

Introns were clustered based on the ratios of intron RPKM to gene RPKM (intron/gene ratio) across all samples (feature vector for clustering = intron/gene ratios at 5-, 10-, 15-, 20-, 60-min 4sU-RNA, total and untagged RNA). Introns were first clustered 100 times using standard k-means clustering (k = 10) and Euclidean distance metrics. Final clusters were then defined as introns always clustered together in the 100 k-means runs. To identify classes among clusters, representative introns were determined first for each cluster as those with the smallest Euclidian distance to the cluster median. Exon/intron ratios of the representatives were then normalized to the exon/intron ratios in 5-min 4sU-RNA. Cluster representatives were again clustered 100 times with k-means for values of k between 3 and 10. For each k, the most frequent clustering was determined. For the final clustering, the k was chosen that maximized this frequency (in this case, k = 4 with a maximal frequency of ~90%).

Statistical tests

Significance of differences between two distributions was assessed using the Wilcoxon rank sum test. This test is based on the ranks of observations and, thus, does not require any assumptions on the type of distribution (nonparametric test). Significance of enrichment was evaluated using Fisher's exact test, which tests for a significant association between different types of classification. Correction for multiple testing was performed using the method by Benjamini and Hochberg (1995) for control of the false discovery rate (FDR).

Data access

Sequencing data including raw sequencing reads have been submitted to the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE31653.

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