Nucleotide excision repair (excision repair) is the main mechanism used by many organisms to remove cyclobutane pyrimidine dimers (CPDs) that are induced in DNA by UV light (1, 2). The mechanism of excision repair has been investigated in considerable detail in several model organisms including Escherichia coli, Saccharomyces cerevisiae, mice, humans, and more recently Arabidopsis thaliana (3, 4). In contrast, although Drosophila melanogaster is an extensively used model organism that has contributed significantly to the development of modern views on ionizing radiation and chemical mutagenesis and to the development of double-strand break repair and recombination models (5), there is rather limited information on nucleotide excision repair in this organism.

Work on excision repair in Drosophila is of special interest because, of all of the model organisms studied, ranging from E. coli to humans, it is the only one that has been reported to lack transcription-coupled repair (TCR) (6–10). This conclusion was further supported by genomic analyses that revealed that the CSA and CSB proteins, which are necessary for TCR (11) and are present in organisms ranging from yeast (Rad28 and Rad26, respectively) to humans, are missing in Drosophila and other insects in the order Diptera (5, 12, 13). Considering the importance of TCR in mutation avoidance (14), the apparent absence of TCR in Drosophila is rather surprising. Therefore, we decided to investigate this phenomenon with new biochemical tools that have become available since the previous experimental studies on the subject nearly 3 decades ago.

In transcription-coupled repair, damage in the template (transcribed strand (TS)), but not the nontranscribed strand (NTS), causes arrest of RNA polymerase (RNAP). The arrested RNAP is recognized by Mfd in E. coli (15, 16) and by CSB in eukaryotes (17). The bacterial TCR process is well-characterized and involves concerted removal of RNAP by the Mfd translocase and Mfd-mediated delivery of repair proteins, resulting in repair at an accelerated rate compared with repair of the NTS (15). In contrast, TCR is not well-understood in higher organisms. The mechanism is unlike that in E. coli, because the CSB translocase does not remove the stalled RNAP (18). Nevertheless, CSB, along with CSA, a WD repeat–containing protein with ubiquitin ligase activity (19), somehow enables the eukaryotic basal repair proteins (which are not homologous to proteins in E. coli) to produce a commonly 3–10-fold accelerated rate of TS repair, which varies depending upon the level of transcription (14, 16, 20).

In the classic TCR assay (14), UV-irradiated cells are incubated for various times to allow repair and then harvested. Genomic DNA is isolated and digested with appropriate restriction enzymes, and then duplicate aliquots from each time point are incubated with or without T4 endonuclease V, which incises sites of CPDs. Following separation of the DNA on a denaturing agarose gel, the gel is probed with gene-specific and strand-specific probes. The level of damage at each time point is calculated from the decrease in full-length fragment caused by T4 endonuclease. The loss of damage with time represents the rate of TS repair, which varies depending upon the level of transcription (14, 16, 20).

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5 The abbreviations used are: CPD, cyclobutane pyrimidine dimer; TCR, transcription-coupled repair; XR-Seq, excision repair sequencing; TS, transcribed strand or template strand; NTS, nontranscribed strand or coding strand; NGS, next-generation sequencing; TSS, transcription start site; TES, transcription end site; CSA and CSB, Cockayne Syndrome group A and B, respectively; RPKM, reads per kbp per million total reads; RNAP, RNA polymerase; nt, nucleotide(s); UVSSA, UV-stimulated scaffold protein A.

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low-resolution and has the potential of missing TCR because it is an indirect method that is based on subtracting two large numbers from one another and because it is limited to analyzing one gene at a time. In contrast, the recently developed XR-Seq (excision repair-sequencing) assay directly captures and identifies the excised oligomers, and thus it directly and simultaneously measures repair throughout the genome (4, 20). Using XR-Seq, we find that Drosophila is proficient at TCR despite the previous reports and the absence of CSA and CSB in this organism.

Results

Excision by dual incision in Drosophila

Drosophila S2 cells were irradiated with 20 J/m² of 254-nm UV light and incubated for various times at 27 °C. Then cells were lysed, and the low-molecular weight DNA was separated from genomic DNA by the Hirt procedure (4). After immunoprecipitation with anti-CPD antibodies, the samples were mixed with an internal control 50-mer DNA oligonucleotide, 3′-end–labeled, and separated on a sequencing gel along with size markers. The excised oligomers are 24–32 nt in length, as is the case in human excision repair. Size distribution of excised oligonucleotides as determined by XR-Seq analysis. A range of 24–32 and median of 27–28 nt was found. Using XR-Seq, we find that Drosophila is proficient at TCR despite the previous reports and the absence of CSA and CSB in this organism.

Processing of the excised oligomers

In humans (20, 21), yeast (22), and Arabidopsis (23, 24), the CPD-containing oligomers are rapidly processed by exonuclease(s) such that the primary excision product has a half-life of about 1 h. As repair progresses, two populations of excision products, one ranging from 24 to 32 nt and centering at 26–27 nt and a second in the range of 15–20 nt and centering at 17–18 nt, are observed. These products as produced in NHF1 cells are illustrated in Fig. 2A. The 15–20-nt degradation products can be seen even at the earliest time point of 30 min and become prominent at 2 h. Thereafter, both the primary and secondary excision products detectable by radiolabeling decline, presumably due to low recovery and labeling efficiency of smaller degraded CPD-containing oligonucleotides (not shown). A comparison of the fate of excised oligomers in human and Drosophila cells (Fig. 2B) reveals an interesting contrast; in S2 cells, no degradation products are detectable in the first 2 h and, in fact, up to at least 12 h (data not shown). This indicates that Drosophila either lacks a nuclease that is present in organisms.
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ranging from yeast to Arabidopsis to humans and that generates a similar degradation pattern in all of these other species, or this degradation intermediate is not stabilized in Drosophila as it is in other organisms.

Transcription-coupled repair in Drosophila

Previous work using strand-specific Southern hybridization did not detect a difference in the rates of repair of TS and NTS in individual Drosophila genes (6–8), and it was concluded that Drosophila does not perform TCR. This notion was further strengthened by genome analyses, which revealed that Drosophila and other Dipteran insects lacked CSA and CSB (5, 12, 13) and UVSSA (25) orthologs. However, it was also pointed out that in the absence of TCR, RNAPII stalled at damage sites in the TS would be expected to interfere with excision repair, as has been shown in E. coli (26), resulting in preferential repair of the NTS (5), which was not observed (6–10). To clarify these conflicting observations and theoretical expectations, we analyzed our XR-Seq data from S2 cells to obtain a global view of strand-specific repair of the Drosophila genome.

From our genomic repair maps of Drosophila, we initially focused on the betaTub60D and ade3 genes for TCR analysis because the lack of strand-specific repair in these genes, as determined by Southern blotting, was considered definitive evidence for lack of TCR in Drosophila (6–8). Fig. 3A shows screenshots of repair of the betaTub60D and ade3 genes in Drosophila as well as the human orthologues of these genes (TUBB3 and GART, respectively) in NHF1 and CSB cells, and Fig. 3B shows quantitative analysis of the TS and NTS repair of these genes. As is visually apparent from the screenshots, the TS of betaTub60D is repaired more efficiently than the NTS. A similar result is seen with repair of ade3. The quantitative data for TS and NTS reads in Fig. 3B clearly show that Drosophila performs TCR similar in magnitude to that seen in NHF1 human cells. A gene-by-gene survey of the genomic repair map (not shown) reveals varied levels of TCR in many but not all Drosophila genes, presumably reflecting varied levels or absence of transcription. For comparative purposes, Fig. 3 shows a screenshott and quantitation of repair for a Drosophila gene, CG10348, in which TCR is essentially absent.

Interestingly, in CSB cells (Fig. 3), there is not only a lack of TCR, but a preferential repair of the NTS in both the TUBB3 and GART genes. This result parallels the preferred NTS repair seen in E. coli lacking the TCR factor Mfd (27). This result in CSB cells is consistent with inhibition of repair by the stalled polymerase and merits more systematic investigation.

Genome-wide TCR in Drosophila

To examine TCR in a large sample of Drosophila genes, we analyzed the CPD repair of all nonoverlapping genes over 1 kbp (5,706 genes). The results are shown in Fig. 4A, with results for different repair time points for S2 cells in the top row, and corresponding time points for NHF1 cells (10,100 genes over 5 kbp) in the row below. Each panel illustrates the averaged repair from the transcription start site (TSS) to the transcription end site (TES) for all of the genes considered, plus the average repair 2 kbp upstream and downstream. As is clear from this figure, in both species, TCR across the gene bodies appears to have begun by 10 min, and TS repair is dominant at 30–36 min. The magnitudes of TCR in human and Drosophila cell lines are at comparable levels. At later time points, after the TS is cleared of damage, the preference shifts to the NTS strand. It takes longer to clear TS damage (over 8 h) in human cells, probably due to longer genes in humans. The robust TCR signal exhibited by S2 cells, which lack CS proteins, contrasts starkly with the complete absence of TCR in mutant human CSB cells, which are compared with NHF1 cells in Fig. 4B. In fact, the CSB cells exhibit a trend toward preferential NTS repair, as seen in the individual TUBB3 and GART genes (Fig. 3).

An interesting difference between the human and Drosophila repair profiles is seen in Fig. 4. As reported previously for NHF1 cells (20), and seen here at the 36-min and 1-h time points, there is a switch in strand preference for repair, from TS within the gene body to the NTS immediately upstream of the TSS. This is because of the well-known fact that upstream of the mammalian TSS, there is promoter- and enhancer-specific transcription “antisense” relative to the gene body (28). These “antisense” transcripts are not found in Arabidopsis (24), and our data suggest that they are absent in Drosophila as well. Instead, a dip in repair of both strands is observed immediately upstream of the TSS in Drosophila, presumably because of interference by transcription factors bound immediately upstream of coding sequences.
Discussion

Contrary to the view held for nearly 3 decades, we show that Drosophila performs TCR. In fact, the magnitude of the TS/NTS repair ratio in transcribed genes is comparable to the ratios of corresponding genes in human cells. We ascribe the previous failure to observe TCR in Drosophila partly to the limitation of the T4 endonuclease V digestion/Southern assay used in previous studies. An important additional consideration is the particular time points at which repair was analyzed. The earliest repair time point commonly used in the previous studies was 4 h post-UV treatment (6–10). TCR may be detected in mammalian cells following 4 h of repair (Fig. 4); however, in yeast, which have much smaller genes, TCR is complete within 1 h (22). It appears that a similar rapid kinetic profile in Drosophila, which also has relatively small genes, likely contributes to the discrepancy between our findings and previous studies.

Our finding of TCR in Drosophila raises a new question: analysis of the Drosophila genome and the genomes of other Dipteran insects has failed to reveal CSA, CSB, or UVSSA orthologs, which are known to be essential for TCR in organisms ranging from plants to humans. The E. coli TCR factor Mfd, which has a different evolutionary history than CSB, exhibits considerable sequence homology in the ATPase/translocase domain of the protein. It is possible that another translocase with great sequence divergence from all known TCR translocases couples transcription to repair upon the encounter of RNAP with a transcription-blocking DNA lesion in Drosophila.
ila. In fact, in *Caenorhabditis elegans*, other chromatin-remodeling factors have been suggested to function in place of CSB for TCR during certain developmental stages (29). There is a need for further work to identify such potential transcription-coupled repair factors in *Drosophila*.

A passive mechanism for TCR has been suggested in which the stalled RNAPII reveals to the basal repair factors lesions that would otherwise be concealed by nucleosomes. This mechanism is based partly upon observations of inhibition of repair by nucleosomes, seen *in vitro* (30), and repair of transcription-blocking lesions unimpeded by the presence of the stalled RNAPII, also seen *in vitro* (11, 31–33). This particular model as related to *Drosophila* is attractive in its independence of CS proteins. Further work is needed to clarify the involvement of CS proteins in TCR as well as in the developmental deficiencies associated with Cockayne syndrome (34).

**Experimental procedures**

**Excision and XR-Seq assays**

S2-DGRC cells were obtained from the *Drosophila* Genetic Resources Center. Cells were cultured at 27 °C in Schneider’s medium with heat-inactivated fetal bovine serum at 10% (v/v). For UV treatments, cells were inoculated into R-150 plates and grown to about 25–80% confluence. Medium was gently removed from the semi-adherent monolayer, cells were irradiated with UV-C, and then fresh medium containing sterilized, conditioned medium was added to cells, and cells were incubated at 27 °C. Excision assays employed 20 J m\(^{-2}\) (Fig. 1) or 10 J m\(^{-2}\) (Fig. 2), and XR-Seq assays employed 20 J m\(^{-2}\). Following predetermined repair times, plates were placed on ice, and cooled cells were harvested by scraping and rinsing with ice-cold PBS. Cells were pelleted, transferred to Eppendorf tubes, washed with cold PBS, and resuspended in 320–340 l of cold TE. Samples were processed for excision and XR-Seq assays as described (27), using 5 l of RNase A and proteinase K and using 9.2% sequencing gels. For both assays, samples were immunoprecipitated with anti-CPD antibody and then either radiolabeled (excision assay) or ligated to adapters and processed for sequencing (XR-Seq). Approximately 13% of the input CPD-containing excision products are recovered using this standard excision assay/XR-Seq immunoprecipitation procedure. One plate of S2 cells per repair time point was sufficient for excision or XR-Seq assay. *Drosophila* possesses photolyases, so cells were kept in the dark or under dim yellow illumination from the time of irradiation to the time of addition of NaCl to cell lysates. XR-Seq of NHF1 (normal human fibroblast) cells was performed as described previously (4). XR-Seq analysis of CSB cell repair utilized data deposited in a prior study (20).
Data analysis

At least 6 million unique mapped reads were obtained for each sample. Analysis of sequencing reads and data visualization were as described previously (35). The alignment genome version was dm6_UCSC for S2 cells and hg38_UCSC for human cell lines. For plotting average repair profiles as a unit gene, we chose the genes with length >1 kbp for Drosophila and >5 kbp for Homo sapiens, and the distance between genes was at least 100 bp for Drosophila and 5 kbp for Homo sapiens. With these criteria, the total number of genes selected was 5,706 for Drosophila and 10,100 for Homo sapiens. Each gene was evenly divided into 100 bins from the TSS to the TES, and 2 kbp (25 bins) upstream of TSS, 2 kbp (25 bins) downstream of TES, and for each bin, from first to last, an average value for each of the selected genes was obtained and plotted. The y axis average reads per kbp per million total reads (RPKM) for each bin was plotted with R. The raw data and alignment data have been deposited in the Gene Expression Omnibus under accession numbers GSE76391, GSE67941, and GSE138846.

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