Human genome-wide repair map of DNA damage caused by the cigarette smoke carcinogen benzo[a]pyrene

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Benzo[a]pyrene (BaP), a polycyclic aromatic hydrocarbon, is the major cause of lung cancer. BaP forms covalent DNA adducts after metabolic activation and induces mutations. We have developed a method for capturing oligonucleotides carrying bulky base adducts, including UV-induced cyclobutane pyrimidine dimers (CPDs) and BaP diol epoxide-deoxyguanosine (BPDE-dG), which are removed from the genome by nucleotide excision repair. The isolated oligonucleotides are ligated to adaptors, and after damage-specific immunoprecipitation, the adaptor-ligated oligonucleotides are converted to dsDNA with an appropriate translesion DNA synthesis (TLS) polymerase, followed by PCR amplification and next-generation sequencing (NGS) to generate genome-wide repair maps. We have termed this method translesion excision repair-sequencing (tXR-seq). In contrast to our previously described XR-seq method, tXR-seq does not depend on repair/removal of the damage in the excised oligonucleotides, and thus it is applicable to essentially all DNA damages processed by nucleotide excision repair. Here we present the excision repair maps for CPDs and BPDE-dG adducts generated by tXR-Seq for the human genome. In addition, we report the sequence specificity of BPDE-dG excision repair using tXR-seq.

tXR-seq | nucleotide excision repair | UV | benzo[a]pyrene diol epoxide | lung cancer

Nucleotide excision repair is a versatile repair pathway that removes a variety of DNA damages, including UV- and benzo[a]pyrene (BaP)-induced DNA damages. BaP, a widespread carcinogen, is the major cause of lung cancer (1). It is produced by incomplete combustion of organic materials and converted to the ultimate mutagen, BaP diol epoxide (BPDE), through enzymatic metabolism (2). BPDE preferentially forms bulky covalent DNA adducts at N2 position of guanines and causes mutations if these BPDE-deoxyguanosines (BPDE-dGs) are not efficiently eliminated by nucleotide excision repair (3). Various methods of varying resolutions have been developed for mapping DNA damage and repair genome-wide (4-9). We previously reported a method, termed excision repair-sequencing (XR-seq), for mapping nucleotide excision repair (6). This method has been used to generate excision repair maps for UV-induced cyclobutane pyrimidine dimers (CPDs) and (6-4)pyrimidine-pyrimidone photoproducts [(6-4)PPs], as well as cisplatin and oxaliplatin-induced Pt-d(GpG) diadducts for the human genome and CPDs for the Escherichia coli genome (10-12).

Although the XR-seq method has been quite useful in determining the effects of various factors, such as genetic background, transcription, posttranscriptional histone modification, and chromatin states, on the timing and efficiency of repair (13), the method in its original form requires the reversal of damage in the excised oligonucleotide (26–27 mers in humans and 12–13 mers in *E. coli*) by enzymatic or chemical means before it can be processed for nextgeneration sequencing (NGS) and generation of repair maps. As such, this method has limitations, because it is not possible to reverse, either enzymatically or chemically, most DNA lesions removed from the duplex by nucleotide excision repair, which is a necessary step for PCR amplification and NGS. We have overcome this obstacle by using appropriate translesion DNA synthesis (TLS) polymerases to convert the excised ssDNA oligomer to the dsDNA form and then amplification by PCR, followed by NGS and alignment of the sequences to the genome. Here we present the application of this approach to the mapping of CPD repair and BPDE-dG repair in the human genome.

Results

tXR-seq of CPD and BPDE-dG. To perform tXR-seq, the human lymphocyte cell line GM12878, which has been extensively characterized by the ENCODE project (14), was either exposed to UV (20 J/m² at 254 nm) or treated with (\pm)-*anti*-BPDE (2 μ M). After allowing 4 h or 1 h for repair, respectively, cells were lysed gently, and low molecular weight DNA, consisting mostly of excised 26–27 mers, was separated from chromosomal high molecular weight DNA by centrifugation.

The basic features of sequencing library generation for tXR-seq are shown in Fig. 1. In brief, the excised oligomers are immunoprecipitated with anti-TFIIH (anti-XPB and anti-p62) antibodies, because the excision product is released in a relatively stable complex with TFIIH, which also protects the excision product from degradation by nonspecific nucleases. This is followed by extraction of the excised oligomers and ligation to adaptors and a second immunoprecipitation with either anti-CPD antibody or

Significance

Benzo[a]pyrene (BaP) is a widespread potent carcinogen found in food, coal tar, cigarette smoke, and industrial smoke. Cigarette smoking is the leading cause of lung cancer, and the mutagenesis in smoking-associated lung cancer is determined by multiple factors, including nucleotide excision repair. We have developed a general method for genome-wide mapping of nucleotide excision repair at single-nucleotide resolution and applied it to generate repair maps of UV- and BaP-induced DNA damage in human. Results show a novel sequence specificity of BaP diol epoxide-deoxyguanosine repair. This general method can be used to study repair of all types of DNA damages that undergo nucleotide excision repair.

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Fig. 1. Schematic of sequencing library construction strategy for tXR-seq. The red asterisk represents a DNA lesion processed by nucleotide excision repair. Excised oligomers are precipitated with TFIIH antibodies, extracted, and ligated to adaptors. Then, the adaptor-containing oligomers are precipitated with anti-lesion antibody, the primer with index is annealed and extended by an appropriate TLS DNA polymerase (pink), and the extension product is amplified by PCR to obtain the dsDNA library for NGS.

anti-BPDE antibody. Human DNA polymerases η and κ are used for primer extension in the presence of purified excised oligonucleotides containing either CPD or BPDE-dG, because polymerases η and κ can bypass CPD and BPDE-dG damage, respectively, in an error-free manner (15–17). This is followed by PCR amplification with index primers and NGS on the Illumina HiSeq 2500 platform.

Length Distribution and Nucleotide Frequency Analysis of Excision Products. The PCR products, approximately 145 bp long, can be seen only in the presence of polymerases η or κ , indicating that the resulting dsDNA libraries exclusively contain the sequence information from the previously excised oligonucleotides attributed to efficient lesion bypass (15–17) (Fig. 24). Fig. 2B shows the length distributions of the oligomers obtained by tXRseq and conventional CPD XR-seq in which the CPD is repaired by CPD photolyase. In both CPD tXR-seq and BPDE-dG tXRseq cases, oligomers in the size range of 24–28 nt predominate, with a median size of 26 nt, in agreement with the wellestablished dual incision mode of the human nucleotide excision repair system (18–20). Importantly, the length distribution of the oligomers obtained by tXR-seq for CPD is in remarkable agreement with that obtained by conventional CPD XR-seq, affirming the validity of tXR-seq.

We further analyzed the base sequence distributions along the excised CPD oligomers obtained by tXR-seq and conventional XR-seq (Fig. 2C). As expected, for CPD oligomers isolated by both conventional XR-seq and tXR-seq methods, positions 19-21 from the 5' end were enriched for thymines (Ts), in agreement with the dual incision pattern of the human excision nuclease system (21, 22). In contrast, for the BPDE-dG oligomers, guanines (Gs) are enriched at positions 19-22, consistent with BPDE-dG monoadducts at one of these positions (Fig. 2D). As a control, when randomly chosen oligomers (10 million) of 26 nt from the human genome were analyzed for sequence composition, all four nucleotides were uniformly distributed throughout the length of the oligomers, with $\sim 60\%$ AT and 40% GC at all positions, consistent with Chargaff's rules (23) (Fig. 2D). We also analyzed the frequencies of di-pyrimidines (TT, TC, CT, and CC) at each position along the excised oligomers of 26-29 nt from both tXRseq and conventional XR-seq for CPD (Fig. S1 A and B). In both experiments, there was strong TT enrichment at the fixed position of 6 nt from the 3' end, further confirming the validity of tXR-seq.

Sequence Specificity of BPDE-dG Repair. Several studies have been conducted on the effects of nearest-neighbor and distantneighbor sequences on the repair rate of BPDE-dG. The data have been interpreted within the context of structural deformity and duplex backbone flexibility caused by the adduct in various sequence contexts, and some general rules have been derived (24–27). Although these pioneering studies have been quite useful, some of the rules were derived from the excision rates of the BPDE-dG adduct from in vitro experiments with human cell-free extracts and oligonucleotide duplexes 135 bp in length (28).

The tXR-seq method, which produces at least ~20 million reads in each biological replicate, provides a model-independent means for analyzing the effects of nearest and distant (one base flanking the nearest bases) sequence neighbors on BPDE-dG excision efficiencies. To secure the vast majority of the excised oligonucleotides used in this analysis containing BPDE-dG at the same position, we classified the sequencing reads from tXR-seq based on read lengths. An analysis of single-nucleotide frequencies along reads 26–29 nt in length showed Gs enriched at two fixed positions (Fig. S2). For 26, 27, and 28 oligomers, we selected those with nucleotide is G at position 20, 21, and 22, respectively. We further filtered out those reads containing G within three bases upstream and four bases downstream of the specific G position to eliminate other possible BPDE-dG sites that could possibly interfere with our analysis. We applied the same criteria to the randomly selected 26, 27, and 28 oligomers from human genomic sequences and then calculated the frequencies of dinucleotides and trinucleotides for both excised oligomers and randomly selected oligomers. Those frequencies for excised oligomers were normalized to their respective frequencies for randomly selected oligomers. We plotted the average fold enrichment of nearest- and distantneighbor base sequence context frequencies for the damaged G base that had been excised in 26-28 oligomers (Fig. 3).

Based on the results, we can draw the following conclusions: First, the fold enrichment of CG context was ~ 2.5 , whereas there was no apparent enrichment of the other dinucleotide contexts (Fig. 3.4). Second, there was a significant effect of the nearest and distant neighbors on the frequency of BPDE-dG excision. In trinucleotide frequency analysis, the excision frequency generally followed the "C>T>A" rule at both the 5' and 3' distant-neighbor



Fig. 2. Analyses of length distribution and singlenucleotide frequencies for the excised oligomers obtained from tXR-seg and conventional XR-seg. (A) dsDNA library for tXR-seq analyzed by 10% native polyacrylamide gel electrophoresis. Primer extension products (1%) were amplified by PCR using the indicated cycles. NC, nontemplate control. (B) Length distribution of excised oligomers from CPD and BPDE-dG tXR-seq (~20 million each) and CPD XR-seq 6 (~10 million) after removal of adaptors and duplicate reads. (C) Single-nucleotide frequencies for 26-mers obtained by CPD tXR-seg (~13 million) and conventional CPD XR-seq 6 (~10 million). (D) Single-nucleotide frequencies for the 26-mers from BPDE-dG tXR-seq (~12 million) along with single-nucleotide distribution of randomly selected 26-mers from the human genome (10 million).

positions (Fig. 3 B and C and Table 1). The 3' nearest-neighbor base also followed the same rule, except for AG*, which had an A>C>T order. For G*A and G*T, the 5' nearest-neighbor bases with the second-highest excision frequency were A and T, respectively. We also selected the specific G at positions 19, 20, and 21 for 26, 27, and 28 oligomers, respectively, and analyzed the sequence context effect in the same way (Fig. S3 and Table S1). The results suggest that the excision frequency was similar and also generally followed the C>T>A rule. Finally, these conclusions must be qualified, because they are based on the assumption of no sequence effect on adduct formation. In fact, BPDE preferentially binds at methylated CpG sites over any other NpG sites (29). In view of these reported sequence preferences for damage formation, genome-wide BPDE-dG damage distribution must be experimentally demonstrated by Damage-seq (10) for a definitive generalization of the sequence effect on repair.

Effect of Transcription on BPDE-dG Repair. Transcription stimulates repair on the transcribed strand (TS) (30). As a general rule, this effect is more apparent in highly transcribed genes and for lesions that are poorly recognized by the core nucleotide excision repair machinery. To explore the effect of transcription on

BPDE-dG repair, we first generated both CPD and BPDE-dG repair maps and integrated the total RNA-seq signal across the entire human genome using tXR-seq reads and ENCODE RNAseq data from GM12878 cells (Fig. S4). We chose the TP53 gene to analyze the effect of transcription on BPDE-dG repair for several reasons. First, it is mutated as a driver both in UVinduced and cigarette smoke-induced cancers. Second, it is transcribed at moderately high levels in many cell lines, including the GM12878 cell line used in this study. Finally, previous highresolution repair studies by ligation-mediated PCR (LM-PCR) on short fragments of TP53 may be used as references for tXRseq data (31). Fig. 4 shows the transcription and repair maps of human chromosome 17, which carries the TP53 gene, at resolutions ranging from megabase to single base or dinucleotide in TP53. This figure also shows tXR-seq data for both CPD and BPDE-dG repair in GM12878 cells, as well as two excision products that can be unambiguously assigned to specific TT and G damages, respectively (Fig. 4). As is apparent from the repair data at all resolutions, the TS for both TP53 and the adjacent WRAP53 gene transcribed in opposite direction were repaired at higher efficiency than the nontranscribed strand (NTS). In addition, for



Fig. 3. Sequence specificity of BPDE-dG repair. (A) Average fold enrichment of dinucleotide excision frequencies in 26, 27, and 28 oligomers derived from BPDE-dG tXR-seq. To calculate the fold enrichment, the dinucleotide excision frequencies from 26, 27, and 28 excised oligomers were divided by the dinucleotide frequencies from randomly selected 26, 27, and 28 oligomers in the human genome. "X" stands for the possible A, C and T bases. "G*" represents the BPDE-dG lesion. Error bars indicate SDs. (*B* and C) Same as in *A*, except that the trinucleotide frequencies were used for excised oligomers and random oligomers. All dinucleotide and trinucleotide frequencies were calculated only in those oligomers that met our selection criteria. The figures represent data from two merged biological replicates.

Table 1. Sequence specificity of BPDE-dG repair

5' distant	XG*	3' nearest	5' nearest	G*X	3′ distant
C>T>A	CG*	C>T>A	C>A>T	G*C	C>T>A
C>T>A	AG*	A>C>T	C>A>T	G*A	C>T>A
C>T>A	TG*	C>T>A	C>T>A	G*T	C>A>T

 G^\star represents the BPDE-dG lesion, and X represents the possible A, C, and T bases.

both damage types, there appeared to be hotspots for repair, although deeper sequencing is needed to be able to comment on the cause and significance of these repair hotspots.

In addition, we analyzed CPD and BPDE-dG repair around the transcription start sites (TSS) and transcription end sites (TES) in NHF1 and GM12878 cell lines, respectively (Fig. 5). The trend was in general agreement with the data obtained from the NHF1 cell line for the repair of CPD and (6-4)PP as measured by the conventional XR-seq method. As expected, all of the repair profiles had repair peaks around TSS, in agreement with the documented high RNA polymerase II density and nascent RNA levels in this region (13, 32–34). In addition, as we reported previously (13), a low level of enriched excision repair near TSS was also seen in CSB cell line, which can be linked to the open chromatin structure close to TSS. In fact, the repair ratio at 1 h for BPDE-dG fell somewhere between that for CPD and (6-4)PP, indicating that BPDE-dG is recognized by the core excision repair complex at an affinity between that of (6-4)PP and CPD (Fig. 5*A*).

Effect of Chromatin States on Repair. Based on DNA sequence elements and histone posttranslational modifications, 15 chromatin states have been defined (35). In previous studies, we found that although these states did not affect cisplatin-induced damage formation, they did affect repair efficiencies, particularly in active promoters, enhancers, and transcribed gene bodies (10, 13). We analyzed all 15 chromatin states for CPD, BPDE-dG, and (6-4)PP repair (Fig. 5B). Again, the general pattern observed in tXR-seq was similar to the patterns seen in CPD and (6-4)PP repair obtained by conventional XR-seq. Interestingly, for BPDE-dG and CPD repair at 1 h after UV treatment, both exhibited high repair levels over active chromatin states but had distinct repair levels over TS and NTS around TSS and TES (Fig. 5 A and B). For BPDE-dG repair, there was only a minor increase on TS compared with NTS, whereas there was much higher repair on TS than on NTS in CPD repair. This finding further confirms the unique BPDE-dG repair characteristics.

The foregoing findings not only show that BPDE-dG as a bulky adduct is processed by the cell in a manner similar to other bulky DNA lesions, but also show that tXR-seq, as a simpler and more versatile method, can be used for damages not suitable for analysis by the XR-seq method.

Discussion

The recently developed XR-seq method has been quite useful for mapping nucleotide excision repair of UV and cisplatin and oxaliplatin damage. However, the method in its original form is applicable only to damages that can be either enzymatically or chemically reversed. Here we have used TLS polymerases to circumvent this obstacle and made the method more widely applicable. In doing so, we have also generated a human genome repair map for one of the major carcinogens, BaP. Interestingly, we found that transcription-coupled repair has only a moderate effect on BPDE-dG adduct, because this lesion, like the UV-induced (6-4)PP, is relatively efficiently recognized by the core human nucleotide excision repair machinery. We also note that LM-PCR analysis of *TP53* hotspots for cigarette smoke-related mutagenesis has led to the conclusion that BPDE adducts form preferentially at G residues in codons 157, 248, and 273 of *TP53*,

and that these cancer-causing mutagenic adducts are in the nontranscribed strand and are repaired at slower rates than other BPDE-dG adducts in the same strand in *TP53* gene, in the context of G*TC (157), CG*G (248), and CG*T (273) (31). The sequencing depth in this tXR-seq was insufficient to detect repair at these sites and thus make a meaningful comparison with repair at other sites to confirm these findings. With new improvements in NGS technology, we expect that deeper sequencing will be affordable and will allow us to address the issue of whether carcinogenic mutations at these sites are isolated with high damage frequency combined with poor repair efficiency to give rise to lung cancers.

We note that although the DNA polymerases η and κ that we used in this study are not applicable to all lesions processed by nucleotide excision repair. However, many TLS polymerases have been identified and characterized in recent years (36, 37); thus, for virtually any bulky DNA lesion, it is possible to use an appropriate TLS polymerase or a combination of two TLS polymerases to accomplish translesion synthesis and construct genomic DNA repair maps. In conclusion, the tXR-seq method



CPD + CTTGGTCTCCTCCACCGCTTCTTGT

BPDE + GCACCCTTGGTCTCCTCCACCGCTTCTT

Fig. 4. Screenshots of CPD and BPDE-dG tXR-seq signals for chromosome 17 along with the RNA-seq profile in GM12878 cells. (*A*–*C*) Representative CPD and BPDE-dG repair map at increasing resolution, zooming in on the *TP53* and *WRAP53* genes. The yellow shaded boxes indicate the zoomed-in regions. For the RNA signal, the "+" indicates genes transcribed from left to right, and the "-" indicates genes transcribed from right to left. For the tXR-seq signal, "+" indicates plus-strand DNA (5' to 3' direction) and "-" represents minus-strand DNA (3' to 5' direction). Green and dark-red lollipops indicate the CPD hotspot codons (286 and 151/152) (41) and BPDE-dG hotspot codons (273, 248, and 157) (31), respectively. (*D*) Two representative oligomers in CPD tXR-seq and BPDE-dG tXR-seq excised from the *TP53*. The purple dashed line indicates the excision positions in *TP53* at different scales. Pink TT (chr17: 7,576,999–7,577,000) and blue G (chr17: 7,576,997) indicate the positions of the respective damages in the *TP53*.



Fig. 5. Effect of transcription and chromatin states on CPD, BPDE-dG, and (6-4)PP repair. (A) BPDE-dG repair profile around the transcription start and end sites of 5,025 selected genes at 1 h in GM12878 cells, and CPD and (6-4)PP repair profiles at 1 h in NHF1 cells. The TS/ NTS repair ratio for CPD is higher than that for BPDE-dG. NTS, nontranscribed strand; TS, transcribed strand. (*B*) Effect of chromatin states on BPDE-dG repair at 1 h in GM12878, and on CPD and (6-4)PP repair at 1 h in SHF1 cells. The subclassified strong enhancer and weak enhancer differ in their distance from the TSS. States 4 and 6 are closer to the TSS, but states 5 and 7 are farther from the TSS. The figures represent data from merged two biological replicates.

opens the door to genome-wide mapping repair of all types of DNA damages processed by nucleotide excision repair at single-nucleotide resolution.

Nonetheless, we note that although tXR-seq is a significant improvement over XR-seq, further technical and computational optimizations are needed for rigorous and quantitative analyses of all damages processed by nucleotide excision repair. The translesion synthesis efficiency and accuracy of the TLS polymerases used need to be taken into consideration, as does the neighboring sequence effect on TLS. In the analysis of BPDE-dG adducts, the elimination of sequences of reads containing Gs three bases upstream and four bases downstream of the adducted G residue precludes the analysis of neighboring Gs on excision efficiency of the adducted G. Although these limitations are important, we believe that we will be able to address them in the near future. The translesion synthesis efficiency and neighboring sequence effects for most TLS polymerases are known, and we expect to determine these factors for the TLS polymerases that have not yet been analyzed. Regarding the neighboring G effects on BPDE-dG adducts, we are developing computational tools to predict with precision the adducted G even when there are neighboring Gs, and thus we will be able to analyze the effect of all sequence contexts on the repair of BPDE-dG damage and the damage induced by other agents, such as N-acetoxy-2-acetylaminofluorene, that specifically attack G residues.

Materials and Methods

Antibodies and TLS DNA Polymerases. The following antibodies were used in this study: anti-mouse IgG (sc-2025), anti-rabbit IgG (sc-2027), anti-XPB (sc-293), and anti-p62 (sc-292) from Santa Cruz Biotechnology; rabbit antimouse IgG (ab46540) from Abcam; anti-CPD from Kamiya Biomedical; and anti-BPDE (clone 8E11) from Trevigen. TLS DNA polymerases η and κ were obtained from Enzymax.

Cell Line and Culture Conditions. Human GM12878 cells were cultured in RPMI medium 1640 (no phenol red) with 15% FBS and 2 mM glutamine at 37 °C in a 5% CO₂ humidified chamber.

UV Irradiation and BPDE Treatment. GM12878 cells were grown to ~80% confluence before UV irradiation or (±)-*anti*-BPDE (MRIGlobal) treatment. The UV (20 J/m² at 254 nm) irradiation was performed as described previously (6), except that 20 mL of GM12878 cells were irradiated in RPMI medium 1640 (no phenol red) in a 150-mm Petri dish. BPDE stock solution (4 mM), which was freshly made by dissolving BPDE in DMSO, was added to the GM12878 cell suspension at a final concentration of 2 μ M, and cells were incubated for 1 h at 37 °C in a 5% CO₂ humidified chamber.

tXR-seq Library Preparation and Sequencing. After UV or BPDE treatment, GM12878 cells were incubated for 4 h and 1 h respectively. At appropriate time points, cells were collected by centrifugation and lysed in ice-cold Buffer A (25 mM Hepes pH 7.9, 100 mM KCl, 12 mM MgCl₂, 0.5 mM EDTA, 2 mM DTT, 12.5% glycerol, 0.5% Nonidet P-40) for 10 min, followed by 40 strokes in a glass hand homogenizer. The chromatin fraction was pelleted by centrifugation at 16,800 \times g for 30 min at 4 °C, and the supernatant was subjected to TFIIH immunoprecipitation by adding 2 μ g of anti-XPB, 1 μ g of anti-p62, and 200 µg of RNase A (Sigma-Aldrich; R4642) per 20 mL of original cell culture. After 3–5 h rotation at 4 °C, 15 µL of recombinant protein A/G Plus-agarose (Santa Cruz Biotechnology; sc2003) per 20 mL of original cell culture was added, and the mixture was gently rotated overnight at 4 °C. The excised oligonucleotides were eluted from the recombinant protein A/G Plus-agarose beads by 100 μL of Buffer C (10 mM Tris-Cl pH 7.5, 1 mM EDTA, and 1% SDS) for 15 min at 65 °C after two washes with Buffer A and Buffer B (25 mM Hepes pH 7.9, 100 mM KCl, 12 mM MgCl₂, 0.5 mM EDTA, 2 mM DTT, 12.5% glycerol, and 1% Nonidet P-40), respectively. The eluted excised oligonucleotides were extracted by phenol-chloroform, and the ethanol-precipitated excised oliogos were incubated with 5 μ L of RNase A/T1 mixture (Thermo Fisher Scientific; EN0551) for 1 h at 37 °C. Before adaptor ligation, another round of phenolchloroform extraction was applied, and a G50 filtration column (GE Healthcare) was used to further purify the excised oligonucleotides.

For one adaptor ligation reaction, 1 μ L of 5' adaptor (20 pmol), 2 μ L of 3' adaptor (40 pmol), and 1.2 μ L of 10 × hybridization buffer (20 mM Tris·HCl, pH 7.5, 200 mM NaCl, and 0.2 mM EDTA) were added into the 5.8 μ L of purified excised oligonucleotides, and the reaction mixture was incubated at 60 °C for 10 min, followed by a 5-min incubation at 16 °C. Then 4 μ L of 5 × ligase buffer, 1 μ L of T4 DNA ligase HC (Thermo Fisher Scientific; 15224-041), 1 μ L of 50% PEG8000 (New England BioLabs), and 4 μ L of dH₂O were added into the mixture, and the ligation reaction mixture was incubated overnight at 16 °C. The 5' adaptor and 3' adaptor were the same as described previously (6).

After phenol-chloroform extraction and ethanol precipitation, the excised oligonucleotides with adaptors were boiled for 5 min and then immediately placed in ice water. Either 2 μ L of anti-CPD or 4 μ L of anti-BPDE was pre-incubated with 5 μ L of Protein G Dynabeads (Thermo Fisher Scientific; 1004D), 5 μ L of anti-rabbit Dynabeads (Thermo Fisher Scientific; 11203D) and 4 μ L of 50% rabbit anti-mouse IgG (Abcam; ab46540) for 2–3 h, followed by mixing with 100 μ L of Reaction Buffer (20 mm Tris-Cl at pH 8.0, 2 mm EDTA, 150 mm NaCl, 1% Triton X-100, and 0.5% sodium deoxycholate) containing the denatured excised oligonucleotides with adaptors. The mixtures were rotated at 4 °C overnight. The subsequent washing, elution, phenol-chloroform extraction, and ethanol precipitation steps were performed as described previously (4).

In the primer extension step, human DNA polymerases η and κ were used to bypass CPD and BPDE-dG damage, respectively. The 10 μ L of excised oligonucleotides with adaptors was mixed with 15 μ L of 2× TLS polymerase buffer (50 mM potassium phosphate pH 7.0, 10 mM MgCl₂, 5 mM DTT, 200 μ g/mL BSA, 20% glycerol, and 200 μ M dNTP) and 3 μ L of 10 μ M RPIn primer (where n represents different index sequences). The single-cycle primer extension procedure is as follows: initial denaturing at 98 °C for 3 min; ramping to 65 °C at 0.1 °C/s and holding for 10 min; ramping to 37 °C at 0.1 °C/s

addition of either 2 μL of polymerase η or 1 μL of polymerase κ into the reaction with mixing; and incubation at 37 °C for 30 min. Then the primer extension product was purified by phenol-chloroform extraction and ethanol precipitation.

The dsDNA library was constructed by PCR amplification using Kapa HotStart ReadyMix with RP1 and RPIn primers, and purified from 10% native polyacrylamide gel as described previously (6). In CPD tXR-seq and BPDE-dG tXR-seq, the primer extension products were amplified for 11 and 14 PCR cycles, respectively. All tXR-seq sequencing libraries were sequenced on the Illumina HiSeq 2500 platform.

Read Processing and Genome Alignment. At least 22 million reads were obtained in tXR-seq. All adaptors in reads were trimmed using Trimmomatic, and duplicate reads were removed by the FASTX-Toolkit (38). Reads >50 mer are filtered before further analysis. The processed reads were aligned to the hg19 human female genome using bowtie with the following command options: -x -q-nomaqround-phred33-quals -m 4 -n 2 -e 70 -l 20-best -p 4-seed = 123 -S (39).

Data Visualization. All sequencing data were obtained from two biological replicates of tXR-seq dsDNA libraries. To compare the repair signal, we normalized all count data by the sequencing depth and visualized them in the Integrative Genomics Viewer (40). Sequencing data for both CPD XR-seq and (6-4)PP XR-seq at 1 h in NHF1 cells were obtained from a previously published dataset (6) [Gene Expression Omnibus (GEO) accession no. GSE67941]. The raw data and bigwig tracks for tXR-seq in this study were deposited in the GEO database, https://www.ncbi.nlm.nih.gov/geo/ (accession no. GSE97675).

Encode Data. GM12878 long nonpolyA RNA-seq [University of California Santa Cruz (UCSC) genome browser accession no. wgEncodeEH000148],

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chromatin state datasets for GM12878 (accession no. wgEncodeEH000784), and NHLF (accession no. wgEncodeEH000792) were downloaded from the ENCODE portal (genome.ucsc.edu/ENCODE/).

Length Distribution and Nucleotide Frequencies. We obtained the length distribution and nucleotide frequencies using custom scripts after adaptor trimming and removal of duplicate reads. To normalize nucleotide frequencies in excised oligonucleotides to human whole genome nucleotide frequencies, we used our published undamaged human genomic sequencing dataset (10) (GEO accession no. GSE82213) and selected size-specific genomic sequences at random for further analysis.

Repair Profiles of Strands and Chromatin State Analysis. The transcript coordinates from hg19 reference genome were downloaded from the UCSC genome browser. The transcripts with an expression score of \geq 300 were taken into account. Among those transcripts, those with another transcript in the 6 kb updownstream vicinity and those <15 kb were filtered out. Regions 5 kb upstream and 15 kb downstream from the transcription start and end sites were binned into 100-bp windows.

The dataset of chromatin states for GM12878 mapped on the hg19 reference genome was downloaded from ENCODE. Reads per kilobase per mapped million reads values were computed and plotted. Two biological replicate sets were combined to generate the boxplots. Bedtools software was used to count the reads for each analysis. All quantitative data were analyzed and plotted using R or GraphPad Prism 6 software.

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