Mapping yeast mitotic 5′ resection at base resolution reveals the sequence and positional dependence of nucleases in vivo

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ABSTRACT

Resection of the 5′-terminated strand at DNA double-strand breaks (DSBs) is the critical regulated step in the transition to homologous recombination. Recent studies have described a multi-step model of DSB resection where endonucleolytic cleavage mediated by Mre11 and Sae2 leads to further degradation mediated by redundant pathways catalyzed by Exo1 and Sgs1/Dna2. These models have not been well tested at mitotic DSBs in vivo because most methods used to monitor resection cannot precisely map early cleavage events. Here we report resection monitoring with high-throughput sequencing using molecular identifiers, allowing exact counting of cleaved 5′ ends at base resolution. Mutant strains, including exo1–/–, mre11-H125N and exo1–/– sgs1–/–, revealed a major Mre11-dependent cleavage position 60–70 bp from the DSB end whose exact position depended on local sequence. They further revealed an Exo1-dependent pause point approximately 200 bp from the DSB. Suppressing resection extension in exo1–/– sgs1–/– yeast exposed a footprint of regions where cleavage was restricted within 119 bp of the DSB. These results provide detailed in vivo views of prevailing models of DSB resection and extend them to show the combined influence of sequence specificity and access restrictions on Mre11 and Exo1 nucleases.

INTRODUCTION

Accurate and efficient repair of DNA double-strand breaks (DSBs) is critical to cell survival and proper genomic function (1). Two evolutionarily conserved pathways exist to repair DSBs: non-homologous end joining (NHEJ) and homologous recombination (HR) (2,3). In NHEJ, DSB termini are rapidly bound by the Ku heterodimer, a ring-like protein complex comprised of Yku70 and Yku80 in yeast (4). Ku binding constrains nuclease activity at DSB ends and promotes the association of downstream proteins critical to repair (5). The Mre11–Rad50–Xrs2 (MRX) complex also appears at DSBs nearly instantaneously and further promotes downstream signaling, such as Tel1-mediated checkpoint activation (6,7). MRX also has functions in tethering the two DSB termini and end processing during NHEJ (8,9). In NHEJ, the two DSB ends are ultimately ligated by DNA ligase IV (Dnl4 in yeast) (10,11).
Unlike NHEJ, HR relies on nucleolytic processing of DSB 5’-terminated strands in a mechanism called end resection. This highly regulated process creates an exposed 3’-terminated strand suitable for donor pairing and extension (12). End resection is proposed to begin through a targeted endonucleolytic incision by Mre11 <100 bp away from the DSB end in a reaction that is dependent on Rad50 and Sae2 in yeast (9,13). After this incision, Mre11 uses its 3’-5’ exonuclease activity to degrade DNA back towards the DSB end, destabilizing Ku (14). Long range resection in the 5’-3’ direction occurs through Mre11-mediated recruitment of the exonuclease Exo1 or the helicase Sgs1 in partnership with the endonuclease Dna2 (15,16). The combined activities of Mre11 backtracking toward the DSB end and Exo1 or Sgs1–Dna2-mediated degradation of DNA away from the break comprises the prevailing bidirectional model of resection (17). The resulting ssDNA is ultimately coated by Rad51 to promote HR (18).

While many proteins involved in HR and resection have been identified, there is much to be learned about the precise coordination of the earliest initiating events as they occur in vivo. Prior in vitro studies on which current models are based analyzed Mre11 nuclease activity on Ku-occluded synthetic dsDNA ends and characterized a primary incision ~35 bp from the terminus on a 70 bp DNA substrate (19). Wang et al. also observed a second cleavage event occurring further downstream at 65 bp from the DNA end in addition to the 35 bp incision when the DNA substrate length was increased to 100 bp (20). Such work led to a hypothesis that Mre11 makes stepwise endonucleolytic incisions from DSB termini to initiate resection (21).

Evidence directly supporting such highly precise models of resection is lacking for mitotic DSBs in vivo. Most current methods used to monitor resection in vivo use qPCR- or Southern Blot-based approaches to quantify ssDNA appearance beginning 100s of bp from the DSB end (22,23), outside of the window examined by in vitro studies. These methods are highly informative regarding resection kinetics but lack the resolution needed to understand the exact events during resection initiation, specifically regarding Mre11 and Exo1 nuclease activity and the factors that promote or constrain their patterns of action, including the precise locations of Mre11 incision points. The covalent linkage of Spo11 to meiotic DSB 5’ termini has permitted more detailed pictures of this process but these advantages are not available for mitotic DSBs (24,25).

We developed a novel in vivo resection assay to map mitotic end resection at a site-specific DSB in budding yeast at base pair resolution. High molecular weight DNA was subjected to primer extension and ligation into circles in a manner that allowed exact counts of the molecules terminated at all possible resection positions within ~350 bp of the DSB. Loss of Exo1 resulted in increased signal throughout that window, most notably at a position ~65–66 bp from the DSB end that proved to be Mre11-dependent, in explicit support of prevailing resection models (20). This pattern was not rigidly positioned but instead moved at different DSB ends in a manner that suggested a possible sequence preference, nor was it singular in that less efficient incision points were also observed. A distinct signal peak at a position ~206 bp from the DSB was Exo1-dependent and suggested a pause point in progressive resection. The pattern in exo1 sgs1 mutant yeast exposed a ‘footprint’ of factors restricting cleavage activity near the DSB. Other mutants revealed dynamic activities at the extreme DSB terminus. These studies provide a precise view of the nucleolytic activities at DSB ends in living mitotic cells and of the sequence- and position-dependent factors that constrain them.

**MATERIALS AND METHODS**

**Yeast strains, growth media and DSB allele**

Haploid yeast were grown at 30°C in either rich medium containing 1% yeast extract, 2% peptone and 40 μg/ml adenine (YPA) with either 2% glucose, 2% galactose or 3% glycerol as a carbon source. Yeast strains were isogenic derivatives of BY4741 (26). The base strain for most strains used in this study was YW3104, itself a derivative of YW1405 that has been previously described (27). All gene disruptions and modified alleles were made using a PCR-mediated technique or a URA3 pop-in/pop-out method and confirmed by PCR and sequencing (28). A genotype of all yeast strains is provided as Supplementary Table S1.

Relative to YW1405, YW3104 bears a previously described point mutation, dnl4-K466A, that abrogates activity of the yeast NHEJ ligase (29). A further change at the ILV1 locus was made to introduce the NdeI site and other sequences used in most experiments; its exact content was related to a prior design concept that is not discussed here. Specifically, we inserted a bacterial ori sequence followed downstream by the HIS3-MX6 selection marker at a genomic location 802 bp upstream of the ILV1 HOcs (Supplementary Figure S1). The fourth change in all strains that were subjected to sequencing-based resection monitoring was the addition of a control allele, which was always the last construction step to yield paired strains with and without the control allele. For this purpose, we identified an appropriate native RE site in the genome (e.g. NdeI) and inserted a construct containing homology to ILV1 350 bp upstream of the RE cleavage site with URA3 as a selection marker. This construct lacked the HOcs but was homologous to the portion of ILV1 that is needed for primer extension and library amplification (Supplementary Figure S1).

**Oligonucleotide primers and probes**

Oligonucleotide primers and probes were from IDT. A complete list of all sequences is provided in Supplementary Table S2. All primers bore 5’ hydroxyls. Primer extension primers were made with hand mixed UMIs at an equal percentage of A, C, G, T and a series of 6 different known and fixed time point barcodes. They were designed to anneal to the RE site DBS-distal to the HOcs such that primer extension reaction would copy both the 5’ primer tail (from the genomic DNA 3’ end) and the genomic DNA (from the primer 3’ end). Replicate time courses from the same strain used different extension primers to ensure that the time point barcodes did not create an unanticipated bias.

**DSB induction paradigm**

Strains containing the ILV1 HOcs with and without the control allele were grown separately overnight at 30°C to
saturation in YPA-dextrose media then inoculated to a low OD$_{600}$ in YPA-glycerol for a consecutive night. When the OD$_{600}$ of the YPA-glycerol cultures was in the range of 0.3 to 0.6 to ensure exponential, asynchronous growth, the strain with the control allele was spiked into the other YPA-glycerol culture at a ratio of 1:20 as determined by OD$_{600}$. The T0 time point was taken by rapidly mixing 45 mL of this YPA-glycerol culture into 5 mL 0.5 M EDTA to quickly quench cellular nuclease activities. Galactose was then added to the YPA-glycerol culture at 2% final concentration to induce HO expression and DSB formation. Another time point was taken 35 min after galactose addition in the same fashion as T0. At 45 min after galactose addition the YPA-glycerol + galactose culture was pelleted and the cells resuspended in an equal volume of YPA-dextrose followed by continued shaking at 30°C. Further time points were taken as above at 60, 75 and 90 min after galactose addition (15, 30 and 45 min after the switch to dextrose, respectively). Time course materials used in ddPCR-based resection monitoring were generated by a similar protocol but without the control allele spike-in and only 1.5 mL of cellular extract was taken from a smaller total growing culture volume without EDTA addition. 

Genomic DNA preparation and quality validation

High molecular weight gDNA was obtained using the Thermo Scientific Pierce Yeast DNA Extraction Kit. Yeast cells were thus lysed without glass beads or damaging chemical treatment. Genomic DNA was subjected to 0.8% agarose gel electrophoresis to confirm its high molecular weight and purity (Supplementary Figure S3). Importantly, DNA is less prone to random fragmentation near the end of a DNA fragment, which was true for all stretches of DNA we studied except for the ILV1-CR side of the DSB at time 0 before DSB formation. Those molecules placed the interrogation region in the middle of a large RE fragment, which resulted in notably higher background as compared to the control allele at T0 (Figure 6A). This phenomenon resolved by T35 because the HO DSB now placed the interrogation region at the end of the source DNA molecule.

Resection monitoring by RE-ddPCR

To measure resection flux through the interrogation window of our sequencing method, we used a ddPCR method based on resistance of ssDNA to RE cleavage. That method is described in detail in Lomonaco et al. (30). Briefly, HinfI was used to digest 300 ng gDNA from the smaller 1.5 mL cultures. HinfI cleavage was confirmed by gel electrophoresis. ddPCR measurements using probes targeting a HinfI site 355 bp from the HOcs and an ACT1 control locus revealed when the HinfI site became resistant to cleavage due to resection.

Resection monitoring by high resolution primer extension sequencing

Quantification of DNA amount and the exact control allele ratio. gDNA was quantified and HOcs curves were established using a dual fluorophore ddPCR assay with primers flanking the HOcs (FAM) and within an undamaged reference gene, ACT1 (VIC). The fraction of uncut alleles was directly measured as the fraction of molecules giving signal with the HOcs relative to the ACT1 probes. The true control allele spike-in percentage was further established in the sequencing assay using DNA extracted from the T0 sample, i.e., prior to DSB formation. That assay used paired probes homologous to the control allele (only present in the spiked-in cells) and the HOcs-containing ILV1 locus (present in all cells regardless of whether they had the control allele). Importantly, all further time points were taken from the same culture so the value from the T0 specimen applied to the later time points also.

Restriction enzyme digestion, primer extension and intramolecular ligation. Twenty micrograms of gDNA were digested by 40 units of NdeI (NEB) in Cutsmart Buffer at 37°C for 1 h. The tubes were then incubated at 65°C for 20 min to deactivate the NdeI. The digested DNA was purified using AMPure XP Beads and eluted using water in a volume of 60 μL. The primer extension reaction used 1 unit of Phusion DNA polymerase in the provided buffer and a custom primer containing a fixed 6 nt time point barcode sequence, 12 nt hand mixed UMIs and several single fixed nucleotides used to locate the UMIs (see below). Primer extension and tagging of molecules was done separately in each time point with different fixed 6 bp barcode sequences. The uncycled extension reaction was 98°C for 3 min, 55°C for 3 min, 72°C for 10 min. Twenty microliter of the eluate following restriction enzyme digestion and purification was added to three separate PCR tubes for each time point during this step, with a final volume of 50 μL in every reaction. The extended molecules from each time point were purified using AMPure XP and eluted using water in a final volume of 40 μL. The entire eluate from each time point was diluted into separate overnight intramolecular ligation reactions at room temperature using T4 DNA Ligase (NEB) at a final concentration of 6400 units in 320 μL in every reaction. The extended molecules from each time point were purified using AMPure XP and eluted using water in a final volume of 40 μL. The entire eluate from each time point was diluted into separate overnight intramolecular ligation reactions at room temperature using T4 DNA Ligase (NEB) at a final concentration of 6400 units in 320 μL reaction volume in the provided buffer. The ligase was then inactivated at 65°C for 10 min.

Amplification and sequencing of custom library. Once UMIs had been covalently ligated to gDNA 5’ endpoints and the ligase inactivated, all time point samples from a given culture’s time course were pooled together and purified into 200 μL water using AMPure XP beads. Therefore, any bias due to PCR amplification and sequencing was expected to apply equally to all time points. Specifically, ligation junctions were amplified with primers flanking the HOcs that were tailed with sequences homologous to Illumina P5 (DSB side) and P7 (upstream NdeI side) adapters used in high-throughput sequencing. This reaction used KAPA HiFi HotStart Polymerase with the following conditions: 95°C for 3 min, 17 cycles of 98°C for 20 s, 60°C for 15 s, 72°C for 30 s, and finally 72°C for 1 min. Twenty microliter of the ligated and purified eluate was added to 10 separate PCR tubes, each a final volume of 50 μL, for every pooled strain time course during this amplification. Products were purified on AMPure XP beads and eluted in 125 μL then a custom Illumina ampiclon library was made through PCR using IDT Illumina-compatible i5 and i7 in-
indexed adapters (IDT10.ULD_Adapter pairs #97 to #106), which identified the time course, i.e. source sample culture. The second adapter PCR also used KAPA HiFi with the following conditions: 95°C for 3 min and 8 cycles of 95°C for 30 s, 55°C for 30 s, 72°C for 30 s. Fifty nanograms of the purified previous amplification product was put into this PCR with a final volume of 50 μl for each library. Ten individual custom amplicon libraries, each representing one complete time course series, were submitted to the University of Michigan Advanced Genomics Core where they were balanced, pooled and sequenced on 2.5% of a shared Illumina Novaseq lane in the 2 × 150 format.

**Bioinformatics analysis**

**Code and file availability.** All code and support files required to execute data analysis are available from GitHub at [https://github.com/wilsonte-umich/resection-seq](https://github.com/wilsonte-umich/resection-seq). The pipeline was executed using the q pipeline manager available at [https://github.com/wilsonte-umich-q-pipeline-manager](https://github.com/wilsonte-umich-q-pipeline-manager). Briefly, a q data file establishes values for sample- and server-specific variables, which then invokes repository file “resection-seq/process_sample.q” via command ‘q submit <data-file>.q’. Each DSB end to be profiled required an indexed FASTA file with the sequence of the reference and control alleles, plus Perl and R scripts defining their structures. Files for all DSBs used in this study are provided in repository directory ‘resection-seq/Bazzano_2021/orig references’. Each sample to be analyzed required a row entry in a metadata file that described the experiments and time points, provided as ‘resection-seq/Bazzano_2021/ResectionMasterStrainTable.txt’.

**Read validation and alignment.** Data from the sequencing core consisted of paired end reads that had been demultiplexed per sample using the Illumina indices but that still had a mixture of all time points per sample. Read 2 (from the P7 primer) was only needed for sample demultiplexing and was not used further. Each read 1 from the P5 primer was analyzed to determine if it matched, in read order, (i) an appropriately long leader sequence corresponding to the DSB-distal end of the resected fragment, (ii) a UMI of the appropriate length, (iii) a single fixed A base, (iv) a known 6 bp time point barcode, (v) two fixed TA bases and finally (vi) sequence corresponding to the DSB-proximal or control-proximal DNA that had been ligated to the UMI during circularization. Reads that failed to match this pattern were discarded.

For matching reads, 36 bp of the proximal sequence was extracted and aligned to the DSB and control reference sequences using Bowtie (31) with options ‘-v 3 -k 1 -m 1 –best’, such that only unique alignments with up to three mismatches were accepted (some reads were expected to map twice to the common sequence shared between the DSB and control alleles). Productive alignments were sorted and grouped by all unique combinations of time point, UMI sequence, and alignment position, which acted as the definitive molecule identifier. Metrics consistently confirmed high data quality, including that (i) there were <2% unaligned reads, (ii) reads only aligned to the predicted strand of the reference sequences and (iii) >95% of reads aligned with no mismatches.

**Purging redundant UMIs.** We next accounted for PCR or sequencing errors introduced into UMIs that would lead to over-counting of the number of truly unique source molecules. We used custom code to implement the directional model of Smith et al. (32), which considers a UMI network to have originated from the UMI node with the highest counts, with UMIs added to an existing network if they have at most one mismatch from a UMI already in the network. Script ‘collapse/collapse_UMIs.R’ implements an efficient algorithm for solving such networks that matched our needs, given that we expected very large numbers of unique molecules mapping to a few specific alignment positions. The algorithm exploits 2-bit encoding of all possible UMI sequences and pre-calculated XOR values for all UMI pairs with a Hamming distance of 1.

Our 12 bp UMIs did not always have sufficient diversity to avoid UMI collisions for the HO or RE-cut ends of the DSB allele; for this reason and for speed we disregarded DSB allele positions that had >500 K mapped molecules, which always included the DSB end itself. In contrast, all positions were always subjected to UMI analysis for the control allele, which had been doped into the yeast pool at a nominal fraction of 5% so that UMI collisions were minimal. The result was a reduction in molecule counts per unique combination of time point, UMI sequence, and alignment position to reflect only true source molecules.

**Discarding failed libraries.** To identify poorly performant libraries we wrote repository script “resection-seq/purge_q_report.pl” to read the output of ‘q report -j all <data-file>.q’ and write a table with summary statistics of all samples, provided for all samples as repository file ‘resection-seq/Bazzano_2021/ResectionOutputSummary.xlsx’. It allowed us to establish criteria that a time point library needed to have at least 3.5 M cell equivalents of input DNA, as determined by the corrected control molecule count, and a background breakage frequency less than 0.03% in the control allele. Libraries failing these criteria were written to repository file ‘resection-seq/Bazzano_2021/BadLibraryTable.txt’. Because time point libraries were initially handled separately, a sample could have many successful time points even if one failed.

**Assembly, normalization and visualization.** For each sample we cross-tabulated read counts such that mapped positions were in rows and time points in columns and wrote two R Shiny web tools to analyze these tables, provided in repository directory ‘resection-seq/server’; ‘visualize’ creates scatterplots for a series of samples as a function of allele position whereas ‘heatmap’ creates plots of position by time point with values expressed by color intensity.

For each tool, the first normalization step was to sum the count of all unique control allele molecules, whether intact or fragmented, to establish the total number of control alleles (and thus haploid cell equivalents) present in the sequenced DNA. We next calculated the inferred number of DSB alleles present in the sequenced DNA by cor-
recting that control allele count by the fraction of control allele-bearing cells in the sequenced mixture, determined by ddPCR from the time 0 specimen as described above. The fraction of all input alleles that mapped to each possible DSB or control allele position was calculated by dividing the molecule count at that position by the actual or inferred allele counts established above. Although extensive resection of the DSB allele could lead to lower total recovered counts at later time points, the calculated fractions remained accurate because they were normalized to the control allele, which was unaffected by DSB processing.

The signal for the control allele was mostly at the NdeI restriction site (Supplementary Figure S4) but the signal at other control positions provided a means of assessing the extent and nature of random fragmentation that occurred in each DNA prep. This pattern included evidence that smaller PCR fragments were sometimes more highly represented in final libraries, so we characterized the background fragmentation by fitting a linear regression to the non-NdeI positions of the control allele. The fraction predicted by that regression was then subtracted from the fraction values at all DSB positions to establish our final normalized values for DSB allele positions, referred to as ‘signal above control’ or ‘DSB – control’. Examination of samples at time 0 consistently validated that the control allele regression line was an excellent model for background fragmentation of the DSB allele (Supplementary Figure S4).

Finally, normalized values were averaged for each time point over all samples of the same culture type (e.g. WT). Heatmaps applied a linear color transformation to the averaged DSB signal above control; the scale of all plots is provided in the figures. The value can be negative due to random fluctuations but is never expected to have large negative values.

Logo plots. Logo plots were generated using http://weblogo.threeplusone.com/. ILV1-PR and CR % GC content (36 and 38%, respectively), were taken into account when generating these plots.

RESULTS

Mapping end resection intermediates in NHEJ-deficient yeast

The DSB system was previously described and includes (i) the HO endonuclease coding sequence placed under control of the native GAL1 promoter and (ii) a single HO cut site (HOcs) in a nucleosome-free region of the ILV1 promoter (Supplementary Figure S1A) (27,33). To facilitate resection monitoring without competing NHEJ, we added a dnl4Δ-K466A mutation to all strains that renders Dnl4 catalytically inactive but able to bind normally to DSBs (29), henceforth referred to as wild-type (WT) with respect to DSB resection. These alleles result in rapid and irreversible generation of a site-specific DSB at ILV1 in >90% of cells within 30 minutes of adding galactose to growing asynchronous yeast cultures (Figure 1A, Supplementary Figure S2). At 45 min after DSB induction, cultures were resuspended in dextrose media to suppress HO expression and stimulate resection (27,30) as measured in Figure 1B using a ddPCR method based on the resistance of resected ssDNA to digestion by a restriction endonuclease (RE) (23). Care was taken to extract high molecular weight genomic DNA (gDNA) given that sheared gDNA is indistinguishable from resection intermediates (Supplementary Figure S3).

We further added a control allele to our strains that lacked a DSB but had (i) a shared common sequence with the DSB allele and (ii) a unique sequence abutting an NdeI restriction site, the same site as found 84 bp distal to the ILV1 HOcs as well as within the shared common sequence (Figure 1C, panel 1, Supplementary Figure S1B, C). Purified gDNA was digested with NdeI followed by annealing and extension of a primer that matched the NdeI site in the shared common sequence, which incorporated a unique molecular identifier (UMI) and a time point barcode into each gDNA molecule (Figure 1C, panel 2, Supplementary Figure S1B-D). Intramolecular ligation created a covalent bond between the UMIs and the 5’ ends of both DSB and control DNA molecules, which might have been created by NdeI, HO, 5’ resection or random shearing (Figure 1C, panel 3). P5 and P7 sequences for high throughput sequencing were added by PCR amplification of pooled ligation products to minimize batch effects (Figure 1D). Because the primer extension primer had a 5’ hydroxyl, only gDNA strands with 5’ phosphates could give final products.

After sequencing, unique source molecules were computationally assigned to the mappable positions of the DSB and control alleles (see Methods). An average of 21M (range 8M to 34M) unique source molecules were obtained per time point (Supplementary File S1). We could not reliably quantify molecules matching the DSB HO and NdeI cleavage positions due to UMIsaturation but could quantify the control locus because we used a mixture in which only 5% of cells carried the control allele. Both loci had a minimal background signal at internal positions before the DSB was formed (Supplementary Figure S4). At 35 min, signal at the DSB allele shifted to the HOcs position, referred to as position 0 (not shown), and at further time points we observed resection signal above background at positions internal to only the DSB allele (Supplementary Figure S4), referred to with negative numbers to indicate how many bases had been removed from the HO cut end. For visualization, we subtracted the control allele background signal from the DSB allele signal and constructed heat maps (Figure 1E). Importantly, normalization to the control allele ensured that all signals reflect the absolute, not relative, fraction of DSB alleles that terminated at each position.

Signal accumulations are evident at distinct positions even in wild-type yeast

Even though resection initiation through our ~350 bp measurement window is expected to be efficient in our WT strain (Figure 1B), we observed intriguing signal patterns that are explored below. Molecules corresponding to the first few base positions near the HOcs progressively increased from 60 to 90 min after DSB induction (Figure 1E, F). Since the DSB had formed by 35 min, this increasing signal at 60 min must be due to end processing associated with attempted NHEJ or resection initiation. There was also accumulation of signal only in the resecting time points at a single −206 position (Figure 1E). Finally, when
Figure 1. Description and validation of resection sequencing by primer extension and circularization. (A) *ILV1* DSB formation efficiency as determined by ddPCR after galactose addition at time 0 to induce HO expression. (B) ddPCR-based resection assay to determine the percentage of ssDNA after DSB induction at a site 355 bp away from the HOcs. Cells were washed into glucose at 45 minutes (arrow). (C) Steps in resection sequencing to map 5′ terminated intermediates at base pair resolution using primer extension to add UMIs, followed by intramolecular circularization and PCR of junctions. (D) Library products were sequenced such that an indexed P5 primer read, in order, the UMI, a fixed time point barcode and genomic bases from the *ILV1* DSB or a control locus. (E) Heatmap depicting the fraction of unique molecules with 5′ endpoints at each position on one side of the *ILV1* HOcs, expressed as an absolute normalized signal above the background fraction of reads aligning to the control locus at similar fragment sizes (see Methods). The NHEJ-defective Dnl4 catalytic point mutant dnl4-K466A is considered WT with respect to resection. (F) Bargraph showing the percentage of unique *ILV1* molecules with 5′ ends within the first 10 bp away from the HOcs for each time point in WT. Negative position designations indicate the number of bases removed from the 5′ terminated strand. Bars are the average ± standard deviation of four biological replicates. (G) Expansion of the region −30 to −90 bp from the HOcs in WT. The color scale is 2-fold more sensitive than (E).

we looked near the −30 and −60 positions suggested as Mre11 cleavage positions by the biochemical experiments of Cejka and Sung, we noted a slight increase in signal at selected positions, especially at −65/−66 (Figure 1E). This signal was weak but compelling due to its location, similar time course as above, and the negative footprint surrounding it where little cleavage was observed (Figure 1G).

Exo1 loss shifts the DSB end resection pattern more than the loss of Sgs1

To add information to the signal patterns above we repeated resection sequencing in strains lacking Exo1 or Sgs1, which are required for the two main long-range resection mechanisms (16,34). While the *sgs1Δ* resection profile looked largely like that of WT (Figure 2A), *exo1Δ* first led to a decrease in reads aligning near the HOcs, especially at the −2 position, consistent with Exo1 itself being responsible for removing a limited number of terminal bases (Figure 2A-B) (35). In marked contrast, we observed a dramatic increase in reads mapping to the −65/−66 bp position in the absence of Exo1 (Figure 2A, C), indicating that Exo1 did not create this signal but instead was responsible for suppressing it in WT cells. There was minimal signal in *exo1Δ* after the −65/−66 bp position until −119 bp from the DSB, after which we observed a steady increase in intermediates throughout the sequencing window. Despite this general increase in reads mapping further away from the DSB terminus in *exo1Δ*, loss of Exo1 led to a specific reduction in read accumulation at the −206 bp position, identifying it as a likely pause point for progressive Exo1-mediated resection (Figure 2A, D). The large difference when comparing *sgs1Δ* and *exo1Δ* resection patterns suggests that Exo1-mediated resection is the preferred mode of end resection initiation near the DSB. Importantly, *ILV1* read accumulations were nearly identical in independent biological replicates (Supplementary Figure S5).

Mre11 is responsible for the peaks at the −65/−66 and other positions

Results above demonstrate that a protein other than Exo1 must create the signal peak at the −65/−66 position, which prevailing models predict should be Mre11. To explore this, we first examined strains with a complete deletion of *MRE11*. Sensitive examination surrounding the −65/−66
position similar to Figure 1G revealed that loss of Mre11 erased the small signal peak observed in WT cells, although intriguingly the negative footprint surrounding that position remained even without Mre11 present (Figure 3A). Here, it is important to note that our method can only map a single 5' terminal position of a source molecule relative to the DSB-distal primer. Thus, we cannot definitively judge how effectively the more DSB-proximal resection intermediates are cleaved relative to the efficient DSB-distal positions. Importantly, mre11Δ also reduced most signal in our interrogation window after the first ~10 bp, including the ~206 peak (Figure 3B).

To establish the actions of Mre11 more definitively we added the mre11Δ-H125N mutation, which abolishes the Mre11 nuclease activity (36), to our WT and exo1Δ strains. The mre11Δ-H125N strain by itself behaved similarly to mre11Δ implicating the Mre11 nuclease as being required for the increase in distal resection intermediates (Figure 3A). Examination of mre11Δ-H125N exo1Δ relative to exo1Δ revealed a complete loss of signal at ~65/−66 and substantially reduced signal at positions −119 and beyond (Figure 3C, D). These results expose an additional stepwise cleavage event mediated by Mre11 at position ~119, 53 bp from position −66. Notably, at both the −65/−66 and −119 positions the signal peaked in EXO1 wild-type strains at 60 minutes and then decreased whereas it remained persistent in exo1Δ mutant strains (Supplementary Figure S6), consistent with a kinetic model in which Exo1 acts later at the activated nick created by Mre11. The −119 position appears to be a major transition point as reads are suppressed in the region between −65 and −119 and increased after −119 through the end of our sequencing window in exo1Δ mutant yeast.

In contrast, we observed a marked increase in intermediates mapping near the HOcs in the mre11Δ-H125N exo1Δ double mutant (Figure 3C, E). Resection mediated by Sgs1-Dna2 is the primary pathway in the absence of both Mre11 nuclease activity and Exo1 (16). Therefore, the increase in reads near the DSB end in mre11Δ-H125N exo1Δ might reflect a shift to resection carried out from the end by Sgs1-Dna2 as opposed to resection beginning from the nick created by Mre11. Intriguingly, analysis of resection kinetics through our sequencing window via the RE-ddPCR assay showed only a small resection defect in mre11Δ-H125N exo1Δ double mutant yeast compared to mre11Δ-H125N and exo1Δ single mutants alone (Figure 3F). There is strong evidence that Mre11 has end-chewing activity that aids in Sgs1 loading (37), but the presence of these intermediates in mre11Δ-H125N exo1Δ indicates that another protein also has such activity.

**Combined loss of Sgs1 and Exo1 reveals the profile of Mre11 activity permissible at a DSB**

It is known that Mre11 is responsible for ~300 bp of resection in exo1Δ sgs1Δ double mutant yeast defective in
Mre11 nuclease activity is necessary to form resection intermediates at $-65/-66$ bp. (A) The first three panels show heatmaps of resection signal for WT, mre11Δ and catalytically defective mre11Δ-H125N strains in the first 125 bp from the HOcs. The next two panels plot the difference between the WT and mutant strains to highlight where signal was higher (red) or lower (blue) in WT as compared to mutant. (B) Fractional resection signal at positions $-203$ to $-209$ bp in WT, mre11Δ and mre11Δ-H125N strains at the 75-min time point. (C) Heatmaps of resection signal for exo1Δ and mre11Δ-H125N exo1Δ yeast up to 300 bp from the HOcs. (D, E) Fractional resection signal 75 minutes after DSB induction in WT, exo1Δ and mre11Δ-H125N exo1Δ yeast at positions (D) $-50$ to $-130$ bp and (E) $-1$ to $-20$ bp from the HOcs. (F) ddPCR-based resection analysis of WT and mutant strains at a site 355 bp from the HOcs. Points are the average of two biological replicates.

long range resection (38). Indeed, ddPCR quantification of ssDNA produced from resection in exo1Δ sgs1Δ Δ yeast showed a decrease in resection efficiency in this strain as compared to WT (Figure 4A). Analysis of exo1Δ sgs1Δ Δ yeast in our sequencing assay revealed an increase in signal at similar positions as observed in the single exo1Δ mutant strain (Figure 4B,D, note the scale difference in Figure 4D). The only locations where reads were decreased in exo1Δ sgs1Δ Δ compared to exo1Δ or sgs1Δ single mutants were at the positions immediately proximal to the DSB termini, indicating that end chewing events in resection are dependent on the presence of Sgs1 or Exo1.

Interestingly, the percentage of unique reads at the $-65/-66$ bp position was only modestly elevated in exo1Δ sgs1Δ Δ as compared to exo1Δ (Figure 4C), which contrasts with the large (>10-fold) increase in reads throughout the sequencing window further from the DSB in the double mutant (Figure 4B,D). We observed intermediates mapping to positions throughout our sequencing window without a drop off in signal, which indicates that Mre11 or another protein is resecting past the positions that can be mapped by the assay (344 bp), even when we extended our window to include reads mapped to the common sequence shared with the control allele (Supplementary Figure S7). The massive increase in signal in exo1Δ sgs1Δ Δ clearly exposed the transition point at position $-119$ (Figure 4B, D). DSB-proximal to this point, the presumptive actions of Mre11 are restricted to only a few selected positions that have an imperfect periodicity (Supplementary Figure S6). After $-119$ bp, the resection pattern shifts in both exo1Δ sgs1Δ Δ and exo1Δ to one indicative of a kinetically slower process, since once again cleavage at the more distal positions would prevent detection of simultaneous cleavage at the more proximal positions. It is noteworthy that major peaks occur at positions $-208$ and $-294$ in the double mutant, which may represent further stepwise incisions generated by Mre11.

Mre11 endonucleolytic activity is influenced by DNA sequence preferences

Broadly, three mechanisms might dictate the specific resection profiles seen above: (i) proteins might ‘measure’ cleavage positions relative to the DSB end, (ii) pre-damage factors such as nucleosomes or other bound proteins might constrain repair protein action, or (iii) repair proteins themselves might show a driving sequence dependence. To distinguish these in our system, we made two deletions near the $-65/-66$ position in WT and exo1Δ backgrounds (Figure 5A, C). The deletions each removed three base pairs either 31–33 or 71–73 bp away from the HOcs, i.e. DSB proximal and distal to the $-65/-66$ position, respectively (Supplementary Figure S1B). Strikingly, the DSB-proximal
Figure 4. Exo1- and Sgs1-deficient yeast show greatly enhanced resection intermediates beyond position −119. (A) ddPCR-based resection analysis of WT and mutant strains at a site 355 bp from the HOcs with combinations of exo1 and sgs1 deletions. (B) Heatmaps of resection signal for exo1Δ and exo1Δ sgs1Δ yeast. The uniquely mappable ILV1 sequences end at position −344. (C) Fractional resection signal surrounding the −65/−66 bp position for exo1Δ and exo1Δ sgs1Δ yeast. The same y-axis scaling is used on both graphs. (D) Fractional resection signal for the first 300 bp away from the DSB for each time point in exo1Δ and exo1Δ sgs1Δ yeast. Note the large difference in y-axis scaling due to the accumulation of many more intermediates in the exo1Δ sgs1Δ strain.

deletion led to a −3 bp shift in the position of the Mre11-dependent peak in both WT and exo1Δ cells, meaning that the same underlying sequence was being cleaved by the Mre11 nuclease even though it was now closer to the DSB end (Figure 5B). Somewhat unexpectedly, the deletion distal to the −65/−66 position changed the pattern of peaks at positions −60 to −66, likely due to their proximity to the deletion (Figure 5D). Both deletions further shifted the resection pattern throughout the distal portions of the sequencing window, including the Exo1-dependent −206 bp position in WT cells and the −119 position in exo1Δ (Supplementary Figure S8). Thus, the signal patterns throughout the interrogation window proved to be highly sequence, not distance, dependent.

Sequencing of 5′ endpoints formed from both ends of the same DSB allele

To expand our base of observation we next examined the opposite side of the ILV1 DSB using the same conceptual approach as in Figure 1 (Supplementary Figure S1A). Because this second side of the DSB corresponds to the ILV1 coding region, we refer to it as ILV1-CR and the sequence studied above as ILV1-PR, for ‘promoter’ (Figure 6A). We observed a higher background in WT and exo1Δ cells for ILV1-CR, but a similar overall pattern was observed on either side of the break (Figure 6B). Intriguingly, a number of reads lined up at the ILV1-CR −206 bp position in both WT and exo1Δ cells. Of primary note was that a peak appeared at −70 bp for ILV1-CR and not at −65/−66 bp as for ILV1-PR, consistent with the sequence, not distance, dependence noted above. Analysis of the underlying sequence, oriented similarly with respect to the DSB, revealed a similar palindromic ‘TCT’ motif, with the cytosine being an efficient incision point (Figure 6C, D). Examination of the region around −119 in ILV1-PR exo1Δ mutants revealed read accumulations at −125 and −153, 55 and 83 bp after the −70 position, respectively. To examine the nucleotides contributing to the exo1-dependent rise in resection signal in both sequencing windows, we first set a cut-off value of 10% or 20% of the highest number of unique reads counted at one position in each exo1Δ mutant. We then generated log plots of the DNA sequence surrounding these 5′ terminated endpoints (Figure 6E, Supplementary Figure S9). We observed a slight preference for A or G in the position immediately before the 5′-terminated endpoint in Exo1-deficient cells. Even though this sequence-specific signal was weak, it is noteworthy because it is different than the TCT sequence noted above. However, a C preference on just the ILV1-CR side of the DSB became apparent for the more restricted set of peaks at the 20% threshold (Supplementary Figure S9).
Figure 5. The predominant Mre11-dependent incision shows sequence dependence more than a strict positional dependence. (A) Diagram of a 3 bp deletion in ILV1 located 31–33 bp from the HOcs. (B) Fractional resection signal from positions −20 to −100 in WT and exo1Δ yeast 90 min after DSB induction with and without the 3 bp deletion at positions −31 to −33. Vertical dashed lines in the top plot mark the location of the positions that were deleted in the bottom plot. A vertical blue line indicates the position expected for the peak in the bottom plot if the distance from the DSB end had been maintained after the 3 bp deletion. (C) Diagram of a 3 bp deletion in ILV1 located 71–73 bp from the HOcs. (D) Similar to (B) for the 3 bp deletion located at positions −71 to −73.

NHEJ and other repair factors influence resection initiation near the DSB terminus

To gain further insight into the obstacles that Mre11 faces near the DSB terminus, we deleted the NHEJ genes YKU70 and NEJ1 and analyzed early resection sequencing intermediates. Intriguingly, the signal at position −65/−66 persisted in yku70Δ, indicating that Ku binding to a DSB end is not uniquely responsible for this Mre11-dependent cleavage position (Figure 7A). However, the more proximal −42 bp peak became more prominent in yku70Δ. Moreover, loss of Ku created an interesting change near the DSB wherein WT showed a progressive increase in signal over time whereas yku70Δ had a nearly constant signal (Figure 7B), which might suggest an easier movement of resection inward from the DSB end. More drastic was the decrease in reads aligning near the HOcs in nej1Δ, which may also signify an increase in the kinetics of resection initiation. Restriction-based ddPCR measurements of end resection kinetics confirmed a higher occurrence of ssDNA at a site 355 bp from the HOcs in these mutants compared to WT (Supplementary Figure S10).

Finally, examination of factors involved in supporting cleavage by Mre11 revealed an increase in reads aligning near the HOcs in sae2Δ and tel1Δ (note the scale difference between WT and these mutants in Figure 7B) and the same near absence of reads at −65/−66 bp as in mre11Δ mutants. These results confirm a slower 5′ degradation process from the DSB end in the absence of Mre11 nuclease activity, specifically at the Exo1-dependent −2 position.

DISCUSSION

End resection is a fast process occurring at a rate of 4 kb/h (34). However, HR must overcome multiple obstacles close to the DSB that dictate the successful transition to long-range resection (39). One such obstacle is repair by NHEJ. We exploited catalytically defective Dnl4 to remove this competing outcome while maintaining normal Ku and DNA ligase IV assembly at DSB ends (29). Despite these obstacles, >20% of HO-cut ILV1 alleles were resected past our ≈350 bp sequencing window within 60 min of suffering the DSB in asynchronous yeast cultures. Resection steadily increased to ≈40% of alleles at 90 minutes but was associated with only limited detectable 5′-endpoint patterns in strains that were wild-type for resection-associated genes. Thus, while resection was clearly initiating in the sequenced regions during the experiment, the transition to progressive long-range resection was normally kinetically very fast such that few endpoint detections accumulated in our observation window.

Mutations further revealed important features of the observed patterns, summarized in Figure 7C, where interpretations are guided by the following principles. When a signal peak is observed, we infer that there is a relative delay (not necessarily a block) in processing leading to increased steady-state signal for those intermediates. When a strain alteration causes such a signal to increase then that delay has been accentuated, which implicates the altered protein as being responsible for normal processing away from that 5′-terminated intermediate. In contrast, decreased signal in
response to a strain alteration suggests that the protein factor normally creates the signal; such an intermediate can sometimes be inferred to be the product of a nuclease’s action. However, a caveat is that primer extension only reveals the most DSB-distal 5′ cleavage point on a DNA strand, so that DSB-proximal increases could result from decreased endonucleolytic activity more distally and vice versa.

Unlike our base dnl4-K466A mutation, yku70 and nej1 mutations removed the Ku and Nej1 NHEJ factors thought to be inhibitory to resection (14,40). The greatest effect of these losses was indeed to decrease the signal near the DSB. Thus, Ku and Nej1 might normally impede further processing away from these positions. Loss of Nej1 had the largest impact and reduced signals to near-background levels, consistent with a function of Nej1 in limiting resection away from the DSB and thus promoting endpoint detections near the DSB (41). An alternative interpretation is that the DSB-proximal signals are not resection intermediates but instead reflect processing associated with futile attempts at NHEJ. These are not exclusive concepts and the fact that signal at the prominent -2 position failed to accumulate in Ku mutant yeast suggests that NHEJ factors might normally, but only temporarily, stabilize DSB-proximal intermediates. Notably, signal at the -2 position was uniquely dependent on Exo1, which identifies Exo1 as the nuclease that cleaves some, but not all, of the DSB-proximal 5′ intermediates (42). In contrast to Ku and Nej1, loss of Mre11 or its nuclease activity led to increased signal at the DSB-proximal positions, most profoundly at −2 bp, with a similar result upon loss of the Tel1 and Sae2 proteins that support Mre11 (7,13,43). Here, the best interpretation is that the signal increase is secondary to decreased DSB-distal endonucleolytic cleavage.

Moving inward from the DSB end we encounter the most critical region for resection initiation. Previous in vitro analysis of Mre11 activity on Ku-occluded dsDNA substrates revealed incisions at 35–45 and 55–65 bp from the end on 70 and 100 bp substrates, respectively (19,20), leading to the prevailing model wherein Mre11 creates a nick at a distance enforced by the Ku protein block (44). We observed a pattern of iterative peaks throughout this region consistent with the model and with the inference of a ‘step-wise’ cleavage at somewhat, although not precisely, regularly spaced positions (21). Although subtle, the pattern of multiple ILV1-PR peaks at −42, −65/−66, −83, −105 and −119 bp positions were evident in wild-type yeast and en-

Figure 6. Comparing two sides of the ILV1 DSB reinforces sequence dependence more than a strict positional dependence. (A) Sequencing scheme to capture intermediates resecting into the ILV1 coding region (ILV1-CR; Figures 1–5 examined resection into the ILV1 promoter region, ILV1-PR). See annotated map and sequences in Supplementary Figure S1. (B) Heatmaps of resection signal up to 300 bp from the HOcs in WT and exol Δ strains sequenced from either ILV1-PR or ILV1-CR in separate libraries. The scale for ILV1-CR strains is 2-fold less sensitive due to higher background levels. The ILV1-CR region marked with an asterisk failed to return any aligned reads for technical reasons due to an adjacent T-rich stretch of bases. (C) Fractional resection signal at positions -60 to −70 bp in WT and exol Δ yeast sequenced from the ILV1-PR. The sequence of the 5′ terminated strand is shown on the upper X-axis. (D) Fractional resection signal at positions −65 to −75 in WT and exol Δ yeast sequenced from the ILV1-CR, with the sequence as in (C). (E) Logo plot analyzing sequences ± 5 bp in either direction from signal peaks in exol Δ yeast sequenced from either ILV1-PR or ILV1-CR. A threshold of 10% of the max peak height was used to determine the plotted peaks: ILV1-PR n = 75, ILV1-CR n = 67. See Supplementary Figure S9 for depiction of these thresholds relative to peak intensities.
enhanced upon loss of Exo1. That increase was strongly dependent on the Mre11 nuclease as evidenced by absence of $-65/-66$ and $-119$ signal in an $exol\ mre11$-H125N double mutant. The relative weaknesses of the $-42$ signal could be secondary to cleavage at $-65/-66$, but $-83$ had substantially less activity, establishing that the different cleavage positions are utilized with different efficiencies.

A central question is what determines the pattern of cleavage at mitotic DSBs in vivo. Models cited above imply that Ku and/or Mre11 protein sizes are critical factors (19,20,44), but the pattern could be enforced by other proteins bound prior to damage such as histones or other proteins (45,46) and DNA repair proteins can have sequence-specific properties (47,48). We addressed these issues using 3 bp deletions in the $ILV1$-PR sequence. Strikingly, the $-65/-66$ peak moved closer to the DSB with the DSB-proximal deletion and thus tracked the local sequence, not the distance from the DSB end. Further evidence for sequence specificity comes from the identification of a similar incision position at $-70$ bp on the $ILV1$-CR side of the DSB whose sequence shared 5 out of 7 bases with the $ILV1$-PR $-65/-66$ peak. Critically, this predominant cleavage position did not move in $yku70$ mutant yeast but the $-42$ position did become the most prominent. Thus, Ku itself is apparently not the primary factor leading to the position of the inferred Mre11 incisions but it may impede access to the more DSB-proximal positions.

It is equally important to consider where resection intermediates were not observed, which was especially noticeable in $exol\ sgs1$, a strain deficient in long range resection thought to mainly support cleavage by Mre11 (38). Resection initiation signal with this strain increased dramatically, although the signal at $-65/-66$ was only modestly increased relative to $exol$. Indeed, most positions up to $-119$ bp had limited read counts despite the poor ability to transition to progressive resection. We infer that the first $\sim 120$ bp of a DSB are a unique zone where a specific protein ‘footprint’ protects the DNA from promiscuous activ-
ity and where Mre11-dependent cleavage is preferred. The source of that protection is not entirely clear, but it is not Ku, Sgs1, or Exo1 based on our data. It might be Mre11-Rad50 itself, although this is difficult to prove with resection data alone.

The genesis of signal after −119 bp is intriguing. In yeast with Exo1 and Sgs1 it is dependent on the Mre11 nuclease. This is especially true for the strong and specific -206 position, which we believe results from exonucleolytic extension away from the more proximal incisions. Consistently, the -206 signal was abrogated in exo1 yeast suggesting it as a pause point during extension by Exo1. We do not know the nature of the obstacle that causes Exo1 to pause but one candidate is the 9–1–1 complex based on recent observations (38). Whatever is responsible is again sequence specific as the location of the peak shifted with the DSB-proximal 3 bp deletions. The signal accumulation at nearly every position past -119 bp in exo1 sgs1 yeast is uncertain; it might be a result of more distal, kinetically slower and pathologic cleavage by Mre11 but we were unable to make an exo1 sgs1 mre11-H125N triple mutant to test this idea (16).

A caveat is the possibility for artifacts arising from the use of the HO endonuclease. HO can remain bound to its cleavage product (49) so we cannot rule out that HO itself is partially responsible for the observed cleavage patterns, although all DSB repair actions occur at HO DSBs and we make our measurements long after repair proteins have engaged the DSB. We also considered whether HO might cleave off-target to create some measured endpoints, especially when the DSB was resistant to resection (50). One peak at −294 in exo1 sgs1 yeast had possible sequence similarity to the HOcs and could represent ‘skipping’ of HO to this alternative position (Supplementary Figure S11). Our studies also have a limited ability to determine the extent to which nucleosomes might help drive resection cleavage positions, in part because we used a DSB in a nucleosome-free region of a gene promoter, which may be less generalizable to other chromatin (33). Such a promoter region is not necessarily devoid of all nucleosome binding, including by fragile nucleosomes and nucleosome variants (51), while at the same time the DSB repair response is known to engage in nucleosome repositioning (52). Finally, because we used asynchronous cultures our results represent an average of all cell cycle stages. We performed DNA content analysis of a subset of our strains in a parallel study, which showed some G1 resection even in WT and that even exo1 sgs1 yeast have some arrest in response to the unrepaired DSB (30). Differences in cell cycle distributions in mutants thus do not appear sufficient to account for the large changes in base-resolved resection patterns, which are much more consistent with a direct mechanistic action of proteins known to bind DSBs.

Our results align with studies that used sequencing approaches to study meiotic end resection. Notably, we observed increased intermediates throughout our sequencing window in exo1 cells that mirrors the effect reported by the Keeney group (24,25). Our data also support the observations that Sac2 and Tell promote efficient resection and a decrease in unique reads mapping near the DSB termini (24,25,53). Keeney described evidence of Exo1-dependent pausing at heterochromatin but it is uncertain how closely that phenomenon corresponds to our precise Exo1 pause point (24).

In summary, results here provide a uniquely high-resolution picture of mitotic DSB resection initiation. They support published models in which Ku and Nej1 inhibit exonucleolytic resection from the DSB end in a manner that is overcome by Mre11 incision with a peak activity ~60 to 70 bp from the DSB end but with evidence for multiple incision points that might occur in a stepwise fashion. Subsequent nick extension by predominantly Exo1 is rapid and processive but also subject to pausing (Figure 7C). However, the cleavage pattern is complex, substantially sequence dependent, and influenced by an inferred protective protein complex that is not strongly dependent on Ku. Critical challenges are to extend these findings across many DSBs and to correlate DNA-based results with equally high-resolution protein binding studies in vivo to understand the nature of the early mitotic repair protein complex.

DATA AVAILABILITY

FASTQ files with sequencing reads are available for all samples from the NIH Sequence Read Archive via accession PRJNA703820. Code and additional summary processed data files are available on GitHub: https://github.com/wilsonte-umich/resection-seq.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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