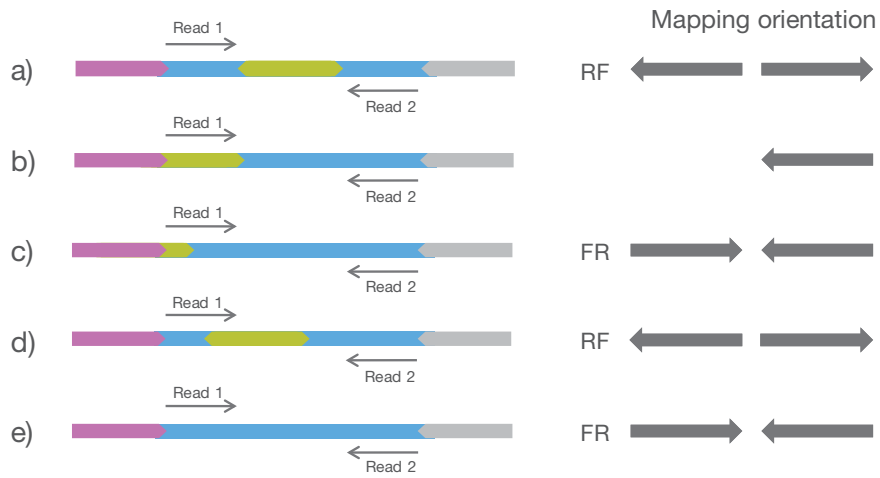




**Figure 2: Junction Adapter Positions and Mapping Orientations**



Composition of example library templates from a mate pair experiment. For each example (a–e), the position of the junction adapter sequence is shown in green and the mapping orientation (either FR or RF, ‘forward–reverse’ and ‘reverse–forward’, respectively) of the resulting read pairs is shown to the right. Sections of genomic DNA sequence are shown in blue and the TruSeq adapter sequences are shown in purple and grey. Amplification/sequencing primer adapters are shown in grey and purple.

**Table 1: Mate Pair Adapter Sequence Elements**

Adapter Element	Sequence
Circularized Duplicate Junction Adapter	CTGTCTCTTATACACATCTAGATGTGTATAAGAGACAG
Circularized Single Junction Adapter	CTGTCTCTTATACACATCT
Circularized Single Junction Adapter Reverse Complement	AGATGTGTATAAGAGACAG
Read 1 External Adapter	GATCGGAAGAGCACACGTCTGAACTCCAGTCAC
Read 2 External Adapter	GATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT

trimming. However, in most cases, the circularized DNA fragments are sheared such that the resulting paired reads will align in the desired RF orientation and will not require junction adapter trimming.

## Adapter Trimming

The junction adapter sequence (Table 1) originates from the Mate Pair Tagment Enzyme used to fragment the input DNA (Figure 1). In most cases, this sequence is appended to both ends of the DNA fragment and therefore is duplicated in an FR orientation during circularization (Table 1, adapter element 1). Less frequently, the adapter is only appended to one end of the DNA fragment, and the post-circularization sequence appears singly (Table 1, adapter elements 2–3).

In addition to these adapter sequences, the templates also contain external adapter sequences used in cluster amplification and hybridization of the sequencing primers (Table 1, adapter elements 4–5). As with standard paired-end library templates, sequencing reads may reach beyond the genomic insert and junction adapter sequence and extend into the external adapter sequence. The likelihood of this occurrence depends on the experimental read length as well as on the length of the final library template.

External adapters must be considered in the alignment process. If they are ignored, the number of reads that successfully align to the reference genome will decrease. For example, a read containing the junction adapter sequence has a very low probability of alignment since that sequence is exceedingly rare in genomic DNA. In cases where an aligner is able to align such a read, the resulting alignment will have a large edit distance, leading to increased mismatch rates and indel error rates.

For these data to be analyzed effectively, the adapter sequences are trimmed prior to alignment. The final orientation of the read pair depends on the location of the adapter sequence within the read and on the applied trimming strategy. Two trim strategies commonly employed are:

1. If the adapter occurs toward the 5' end of the read (Figure 2c), the read sequence after the adapter is kept. Together with its partner this will yield paired-end reads in an FR orientation.
2. If the adapter occurs towards the 3'end of the read (Figure 2d), the read sequence before the adapter is kept. Together with its partner this will yield mate pair reads in an RF orientation.

When correctly oriented mate pairs are sought, it may be preferable to





