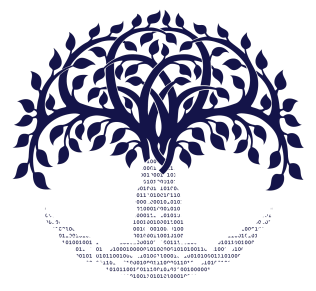
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# DTU Health Technology Bioinformatics

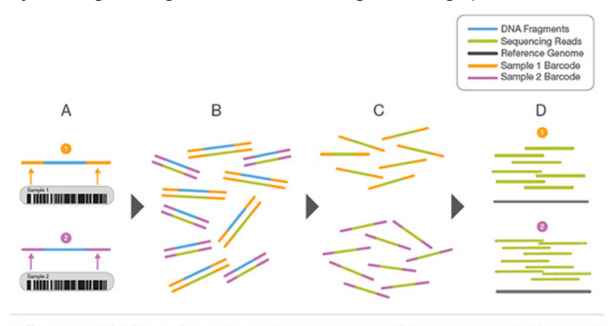
### **Barcoding**

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## Multiplex sequencing

 Multiplex sequencing allows sequencing libraries to be pooled and sequenced simultaneously during a single run on most high-throughput instrument



- A. Two representative DNA fragments from two unique samples, each attached to a specific barcode sequence that identifies the sample from which it originated.
- B. Libraries for each sample are pooled and sequenced in parallel. Each new read contains both the fragment sequence and its sample-identifying barcode.
- C. Barcode sequences are used to de-multiplex, or differentiate reads from each sample.
- D. Each set of reads is aligned to the reference sequence.

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- NovaSeq system outputs 20 billion paired-end reads yielding up to 3000 Gbases
- Running several samples on one sequencing run
  - Saves money & time
  - Eliminates sequencing run as variable

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### **De-multiplexing**

- De-multiplexing is generally done automatically before you receive the samples
- If sequencing locally always check how much on the run ended up unassigned
  - Bcl2fastq2 stringency can be adjusted if barcodes are very different



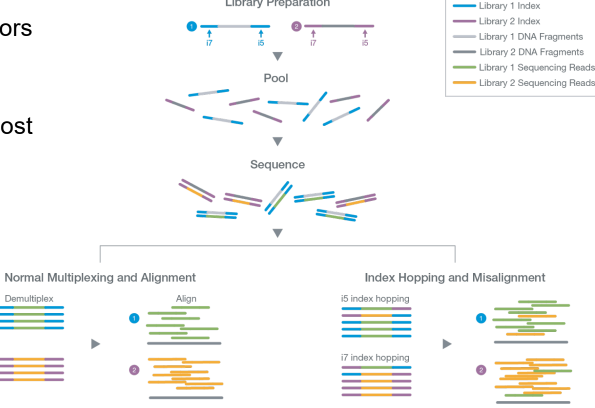
How would you realise that demultiplexing was not done?

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## Index hopping and bleeding

- Failure to remove free adapters leads to index hopping
- Very similar barcodes and sequencing errors can lead to index bleeding
- Dual indexing solves both problems at a cost



**Library Preparation** 

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