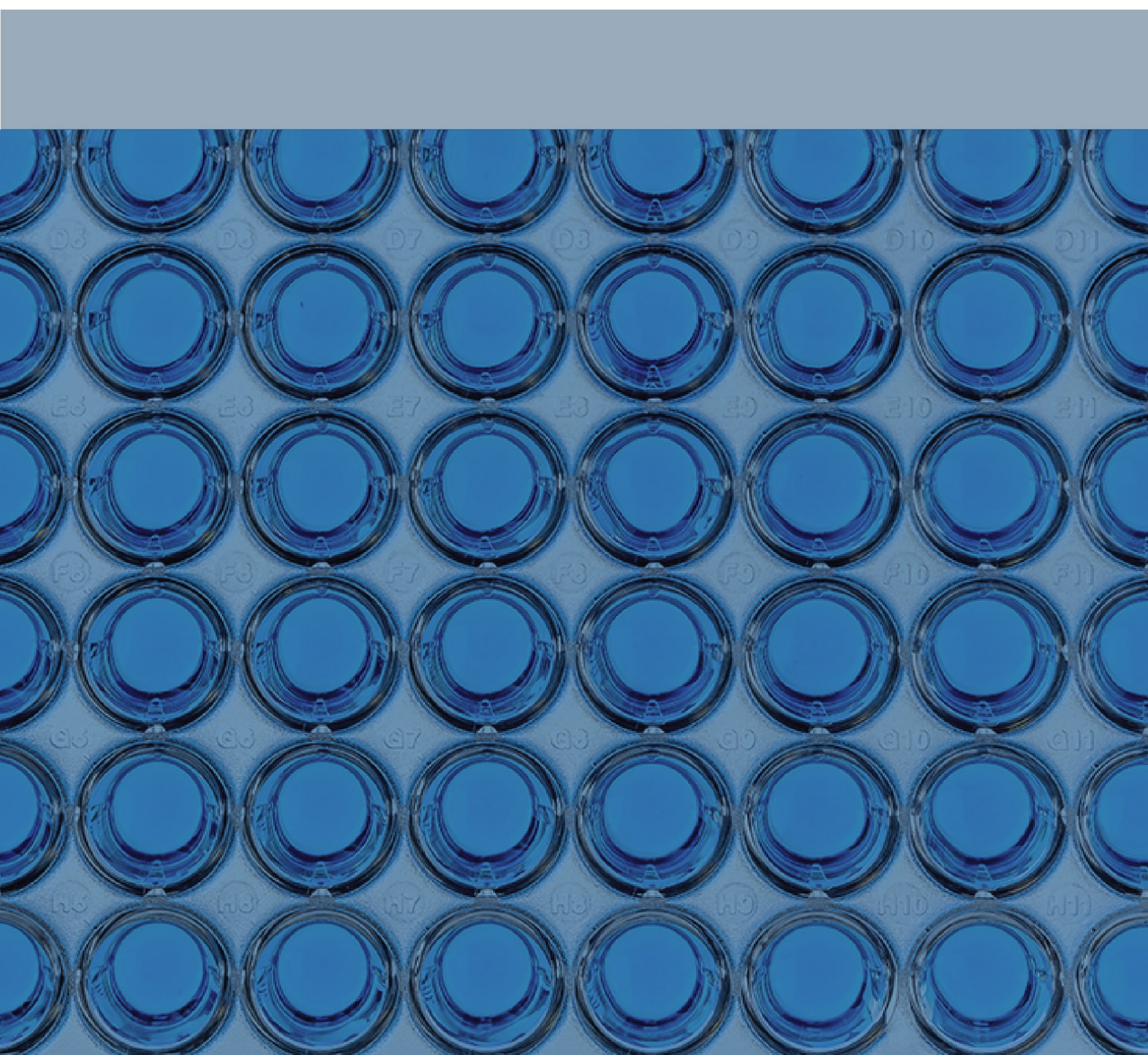


# purePlex™ DNA Library Prep Kit

**Featuring speed, batch flexibility, and data confidence with UDIs.**

seqWell's core technology is a true multiplexing library prep system for making normalized NGS libraries quickly and easily from large numbers of samples.



# purePlex™ DNA Library Prep Kit

Featuring speed, batch flexibility, and data confidence with UDIs.

## Principal Features

## Reduces labor costs and QC burden

Fast, flexible workflow with no requirement for full plate processing

Auto-normalization reduces QC burden, improves data consistency

Early pooling for easier sample handling

Reduced GC bias compared to other transposase-based methods

Accurate sample identification

384 UDIs available (4 sets of 96)

The purePlex DNA Library Prep Kit leverages our patented Tn5 transposase-based chemistry where simple molecular tagging steps add dual indexes early on in the library prep process, enabling pooling of samples prior to any bead cleanups or amplification steps.

A distinct advantage of the purePlex workflow is the elimination of individual sample and library normalization which reduces the QC and labor costs compared to traditional UDI workflows.

## Application Strengths

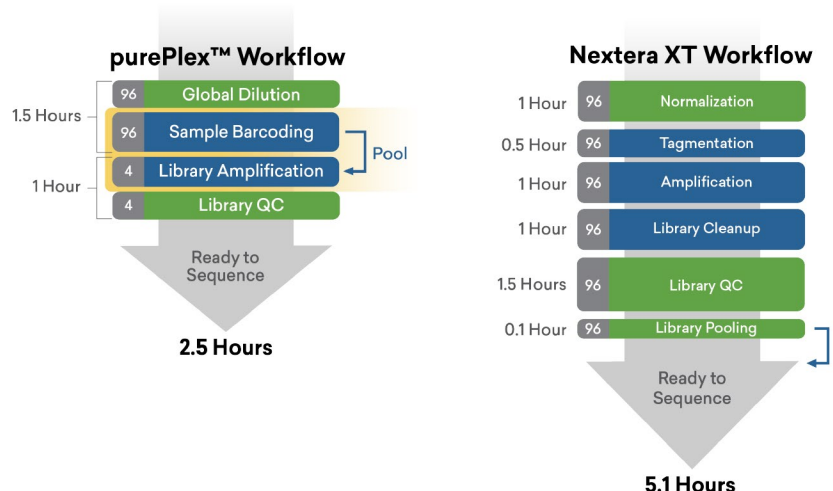
## From samples to loading sequencer in less than a day's work

Low-pass whole genome sequencing

Whole small genome sequencing (<50 Mb)

Metagenomics/Microbiome screening  
scRNA-seq

Synthetic construct sequencing  
(amplicons, plasmids, etc.)



2.5-hour workflow for preparing 96 samples  
45 min. hands-on time



Save money and the planet  
through using less plastic

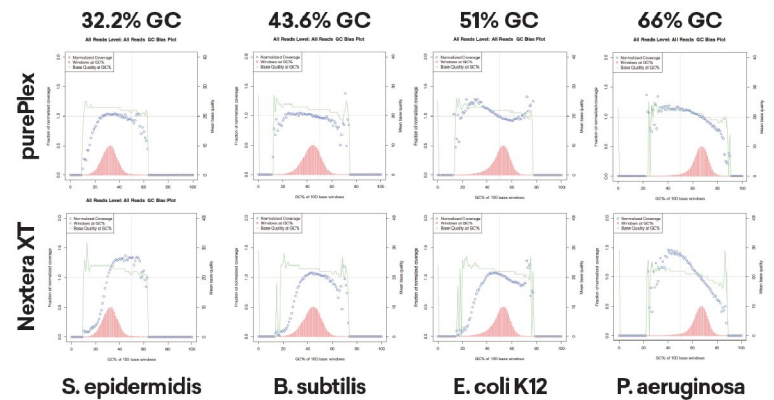


Robust performance for  
all GC contents

GC bias plots demonstrate purePlex has  
more even coverage across high and low  
GC regions compared to Nextera XT.

“With our streamlined workflow, you can save  
more than 50% in plastics per sample compared to  
Nextera XT, based on our estimates. And that results  
in savings of \$7 or more per plate.”

– Maura Costello, Team Leader, R&D, seqWell

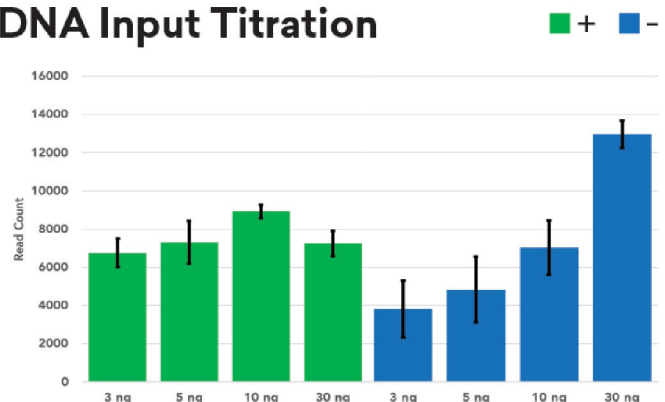


Auto-Normalization of  
Read Depth

Samples were normalized to inputs of 3, 5,  
10, and 30 ng then underwent purePlex  
library prep with (+) and without (–)  
normalization reagent.

Read counts for each sample are equal,  
regardless of input, when normalization  
reagent is used. In contrast, without  
normalization reagent, sample read  
count scales with input.

## DNA Input Titration



## Specifications

### purePlex™ DNA Library Prep Kit

Catalog #301067, #301068, #301069, and #301070

Specifications	Description
Sample Type	Amplicons, Plasmids, Genomic DNA, cDNA
DNA Input Range	5-50 ng
Number of Unique Index Combinations	2304
Supported Paired Reads (Clusters/Sample)	≤ 20 million
Output Fragment Range	400 – 1,200 bp
Primary Applications	Synthetic construct sequencing (amplicons, plasmids, etc.), Low-pass whole genome sequencing, Whole small genome sequencing (<50 Mb), scRNA-seq, Metagenomics/Microbiome screening
Reactions per Kit	96



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