B-cell receptor reconstruction from single-cell RNA-seq with VDJPuzzle

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\textbf{Abstract}

The B-cell receptor (BCR) performs essential functions for the adaptive immune system including recognition of pathogen-derived antigens. Cell-to-cell variability of BCR sequences due to V(D)J recombination and somatic hypermutation (SHM) necessitates single-cell characterization of BCR sequences. Single-cell RNA sequencing (scRNA-seq) presents the opportunity for simultaneous capture of the BCR sequence and transcriptomic signature for a detailed understanding of the dynamics of an immune response.

We developed VDJPuzzle 2.0, a bioinformatic tool that reconstructs productive, full-length B-cell receptor sequences of both heavy and light chains. VDJPuzzle successfully reconstructs BCRs from 98.3\% (n=117) of human and 96.5\% (n=200) from murine B cells. 92.0\% of clonotypes and 90.3\% of mutations were concordant with single-cell Sanger sequencing of the immunoglobulin chains. VDJPuzzle is available at https://bitbucket.org/kirbyvisp/VDJPuzzle2
Introduction

B cells are an essential component of the adaptive immune system. Specificity of B cells for their antigen is imparted by the BCR, which comprises a paired heavy (IGH) and light (IGK or IGL) immunoglobulin chain that differs from cell to cell. Each of these chains contains one of many possible V, J, and constant segments encoded by clusters of genomically-adjacent genes. Additionally, IGH includes a D segment between the V and J segments. The specific combination of genes is selected through genomic rearrangement in a process called V(D)J recombination and occurs during B-cell development [1].

After V(D)J recombination, naïve B cells are activated by binding an antigen through the BCR. Additional BCR diversity and specificity for antigen is generated through a process called affinity maturation during which SHMs accumulate in the germline sequence of the V(D)J genes. Preferential mutation occurs in the complementary-determining regions (CDRs) which directly bind to antigen; the most variable of these is the complementarity determining region 3 (CDR3) which adjoins the V(D)J junction. This region determines the affinity of the BCR for the cognate antigen.

Obtaining high-quality BCR sequences along with single-cell transcriptomics is crucial for characterizing the dynamics of B-cell evolution and differentiation during an immune response. scRNA-seq enables the simultaneous measurement of these important parameters [2, 3]. Similar approaches for reconstruction of full-length T-cell receptors (TCRs) have yielded insights into T cell biology [4-8], including determination of T-cell development trajectories in the tumor microenvironment [9].

Here, we present VDJPuzzle 2.0, a computational method to reconstruct the full length BCRs from scRNA-seq and at the same time also provide gene expression data. This tool extends a software package previously developed to reconstruct T-cell receptors [6]. The new VDJPuzzle version has been adapted for BCRs and includes additional features to reliably characterise SHMs.

Results

Algorithm

For each cell, trimmed paired-end reads are aligned to the reference genome. Pairs with at least one read aligning to any of the V(D)J or constant-region genes are extracted and used to assemble contigs using the de novo assembler Trinity [10]. Assembled contigs are matched to the IMGT database using MiGMAP (https://github.com/mikessh/migmap) to find complete, productive, and in-frame BCRs [11]. VDJPuzzle workflow is represented in Fig. 1.

Success rate and quality of reconstructed BCR sequences

VDJPuzzle was applied to scRNA-seq data from 117 B cells isolated from human peripheral blood mononuclear cells (PBMCs) derived from a healthy donor. These B cells were additionally sorted by surface phenotype: 27 transitional, 30 naïve, 30 memory, and 30 plasmablast B cells. We define a successful BCR reconstruction as a chain with a complete, productive, and in-frame BCR sequence identified using
IgBLAST run via MiGMAP. VDJPuzzle successfully reconstructed at least one V(D)J sequence of a heavy chain (IGH) and at least one light chain (IGL or IGK) in 98.3% (n=115). Of the 115 B cells, 72 (62.6%) contained at least one IGK, while 54 (47.0%) had at least one IGL (Table 1). In 7.0% (n=8) cells, VDJPuzzle reconstructed both IGK and IGL, consistent with previous findings of isotypic inclusion within the normal B cell repertoire of humans and mice [12].

Sanger sequencing from the single-cell cDNA libraries for the 117 B cells was performed as validation of the VDJPuzzle output. After filtering low-quality traces, Sanger sequencing reconstructed BCR transcripts in 95 and 80 cells, respectively. Out of the 95 IGH, 89 had concordant clonotypes (i.e. identical CDR3 and V-J genes) compared to the corresponding VDJPuzzle-reconstructed BCR. Two cells had the same CDR3 sequence, but discordant V and J genes. Two other cells had both different V-J genes and CDR3 sequences. VDJPuzzle failed to reconstruct the receptor in two other cells. A total of 78 light chain (24 IGL and 52 IGK) sequences from the 80 reconstructed with Sanger matched the VDJPuzzle result in both CDR3 and V-J genes. The two discordant chains contained either a single amino acid difference in the CDR3 sequence, or had the same CDR3 but different V-J genes.

Analysis of mutations with respect to the germline from the 89 heavy chains and 78 light chains with concordant clonotypes between Sanger and VDJPuzzle revealed that 2376 (97.7%) mutations in the heavy chain and 1666 (94.9%) in the light chain were shared with Sanger. In contrast, 57 and 89 were identified only by VDJPuzzle, while 141 and 226 mutations were identified by Sanger alone. As further comparison, a recently-published BCR reconstruction method called BASIC was tested on the same dataset [13]. Only 6.0% (n=7) of heavy chains and 21.4% (n=25) of light chains were complete, productive, and in-frame as determined by IgBLAST [14]. However, BASIC success rate may improve with a different dataset or when a different definition of successful reconstructed BCR is used. Indeed, Canzar et al. reported a success rate of 97% in identifying the correct V(D)J gene usage.

As alternative comparison, a less stringent success rate has been utilized to benchmark VDJPuzzle, BASIC [13], and Sanger sequencing. In this analysis, we allowed MiGMAP to report BCR with incomplete, not in-frame, non-coding, and not canonical BCRs. Results are showed in Table 2. Both VDJPuzzle and BASIC improved, however this also increased the number of cells with multiple heavy and light chains due to the inclusion of non-productive BCR.

Next, we examined germline mutations as a function of IMGT-annotated framework (FR) and CDR regions [11]. As expected, more SHMs occurred in the CDR regions compared to the FR regions. Further, plasmablasts and memory cells carried more mutations, while naïve and transitional B cells had a lower number of SHMs (Fig. 2). To demonstrate the utility of the VDJPuzzle output, we linked the aggregate mutation rate across the entire V(D)J region with the gene-expression data from scRNA-seq (Fig. 3). Unsupervised clustering of the transcriptomic data identified markers of naïve/transitional, plasmablast, and memory populations. Known terminal differentiation markers such as XBP1 were highly expressed in the
plasmablast cluster [15], whereas expression of known early differentiation markers including TCL1A was observed in naïve/transitional cells [16]. Furthermore, a cluster of co-expressed genes was negatively correlated with mutation frequencies (R=-0.79, p-value<0.001).

VDJPuzzle was also tested on mouse B cells. Out of 213 cells in ArrayExpress (E-MTAB-4825), 13 were not available for download on the FTP server. From the remaining 200 cells, VDJPuzzle reconstructed 193 heavy chains (96.5%) and 200 light chains (100%). 195 cells (97.5%) had at least one IGK, while 8 (4%) had at least one IGL. 3 cells (1.5%) had both IGK and IGL.

**Discussion**

VDJPuzzle achieves a high success rate in reconstructing BCR sequence from scRNA-seq compared to existing methods. Other software can achieve high detection of V(D)J usage and the CDR3 from RNA-seq data, such as MiXCR [17]; however, these programs cannot reconstruct the full-length BCR sequence and do not integrate these data with gene expression. We anticipate that simultaneous determination of BCR sequences with transcriptomic signatures from single cells will be a promising approach for studying the dynamics of an immune response in heterogeneous populations of B cells.

**Methods**

**Single cell sorting**

PBMCs were separated from freshly collected whole blood by density gradient centrifugation and stained with the following monoclonal antibodies: CD19-BV421 (Biolegend), IgD-PerCP-Cy5.5, CD27-PE-Cy7, CD38-APC, CD10-PECF594, CD3-APC-Cy7, IgG-BV605 (BD Biosciences) and viability dye eFluor780 (eBioscience) for 30 min on ice and, following two washes, cells were resuspended in PBS with 1% HI bovine serum for single cell sorting. Following exclusion of non-viable/CD3+ cells, individual transitional (CD19+, IgD+, CD27-, CD10+), mature naïve (CD19+, IgD+, CD27-, CD10-), switched memory (CD19+, IgD-, CD27+, IgG+) and plasmablast (CD19+, IgD-, CD27++, CD38++) were sorted directly into 96-well PCR plates (Eppendorf) using a FACS ARIA III (BD Biosciences). PCR plates were stored at −80 C for subsequent analysis.

**Single cell RNA-sequencing protocol**

Single cell RNA sequencing was performed as described by Picelli et al. [18, 19] with the following modifications. Cell lysis buffer was prepared by adding 1μL Rnase inhibitor (Clontech, Mountain View, CA) to 19μl Triton X-100 solution (0.2% v/v). Cells were sorted directly into 96-well plates containing 1μL lysis buffer, 0.5μL dNTP (10mM) and 0.5μL oligo-dT primer at 5μM. Reverse-transcription and PCR amplification were performed as described with the following exceptions: reactions were performed at half
volumes, the IS PCR primer was reduced to a 50nM final concentration and the number of PCR cycles increased to 28. Sequencing libraries were prepared using the Nextera XT Library Preparation Kit (Illumina; San Diego, CA, USA) at one quarter of the recommend total volume and sequencing performed on the Illumina NextSeq sequencing platform.

**Single cell RNA-sequencing data analysis**

Reads obtained from scRNA-seq experiments were used as input of VDJPuzzle which perform the BCR assembly (method section of the main text), it aligns the reads to the ensembl reference genome (GRCh37), and it quantifies the gene expression of each cell. VDJPuzzle alignment step utilized TopHat2 [20], while gene expression is calculated with CuffLinks 2.2.1 [21]. Downstream analysis of gene expression has been performed in R. Clustering analysis have been performed with SC3 R package [22].

**Sanger sequencing protocol**

Sanger sequencing was performed fromorm the same cDNA libraries utilized for scRNAseq for the 117 B cells and for both heavy and light chains. Two ng of amplified SmartSeq2 cDNA was subjected to 35 cycles of PCR using forward primers binding leader peptide sequences of immunoglobulin heavy and light chain V segments and reverse primers binding μ, γ, κ, and λ constant regions [23]. PCR products were Sanger sequenced with the same forward and reverse primers by Genewiz (Boston, USA).

**Sanger sequences data analysis**

The 468 tracing files (forward/reverse for each chain and each cell) were loaded on Geneious [24] and manually trimmed based on the chromatogram quality. Forward and reverse sequences were merged together to create a consensus sequence for each chain and each cell. The sequences were loaded into MiGMAP to identify the V(D)J usages, FRs and CDRs regions, along with the mutations compared to the germline sequence of the matched BCR in IMGT.

**Single cell RNA-sequencing of mouse B cells**

Mouse scRNA-seq were downloaded from ArrayExpress (E-MTAB-4825) [3]. Data analysis was performed as for human PBMC using GRCm38 reference genome on the 200 B cells retrieved from the ArrayExpress ftp server.

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*Conflict of Interest: *none declared.
References


### Tables

#### Table 1. Benchmark results of the VDJPuzzle, BASIC, and Sanger sequencing.

<table>
<thead>
<tr>
<th>Method</th>
<th>IGH</th>
<th>IGK</th>
<th>IGL</th>
</tr>
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<tbody>
<tr>
<td>VDJPuzzle</td>
<td>115 (98%)</td>
<td>72 (61.5%)</td>
<td>54 (45.3%)</td>
</tr>
<tr>
<td>BASIC</td>
<td>7 (6.0%)</td>
<td>19 (16.3%)</td>
<td>6 (5.1%)</td>
</tr>
<tr>
<td>Sanger</td>
<td>95 (81.2%)</td>
<td>55 (47%)</td>
<td>25 (21.4%)</td>
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#### Table 2. Alternative benchmarking. Numbers and percentage of reconstructed chains considering only BCRs with at least one V and J (no D and CDR3 required).

<table>
<thead>
<tr>
<th>Method</th>
<th>IGH</th>
<th>IGK</th>
<th>IGL</th>
</tr>
</thead>
<tbody>
<tr>
<td>VDJPuzzle</td>
<td>117 (100%)</td>
<td>104 (88.9%)</td>
<td>103 (88.0%)</td>
</tr>
<tr>
<td>BASIC</td>
<td>59 (50.4%)</td>
<td>59 (50.4%)</td>
<td>55 (47.0%)</td>
</tr>
<tr>
<td>Sanger</td>
<td>96 (82.1%)</td>
<td>55 (47%)</td>
<td>26 (22.2%)</td>
</tr>
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</table>
Fig. 1. VDJPuzzle workflow.
Fig. 2. Mutation rate across FR and CDR regions. Mutations are called comparing the sequence against the BCR germline sequence. The number of mutations is normalized by the length of the FR/CDR regions.
Fig. 3. Clustering analysis of scRNAseq and relationship with SHMs. Unsupervised clustering was performed using SC3. Naive and transient B cells are clustered together and associated to a low mutation rate. In contrast, Memory B cells and Plasmablast form two clusters which are associated to high mutation rate. Notably, the gene signature (gene cluster 1) associated to naïve and transient B cells was negatively correlated to the average mutation rate identified in each cell.