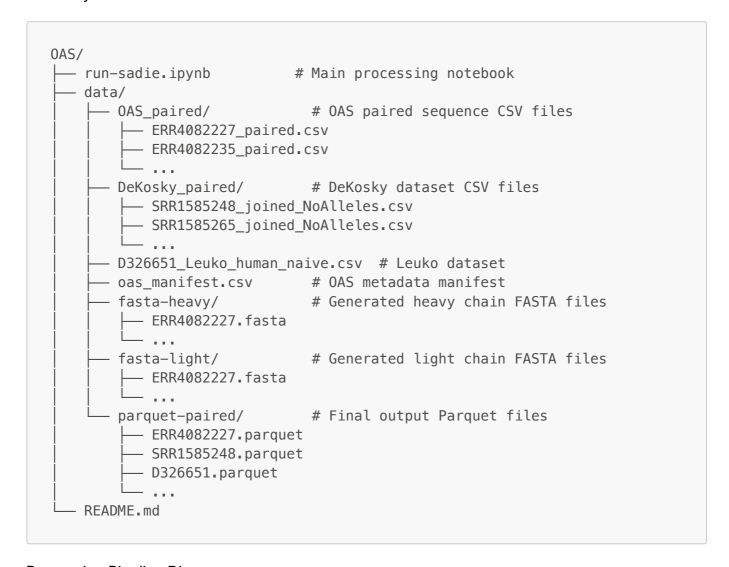
#### Bader2025

This is the public code and data repository for Bader et al. 2025.

#### **Abstract**

This pipeline processes paired heavy and light chain antibody sequences from the Observed Antibody Space (OAS) database, DeKosky and Leuko datasets. The sequences are annotated using SADIE's IgBLAST integration and saved as Parquet files with complete AIRR-compliant annotations and metadata for downstream analysis.

# **Directory Structure**



# **Processing Pipeline Diagram**

INPUT DATA SOURCES		
OAS Database (CSV files)	DeKosky Data (CSV files)	Leuko Dataset (Single CSV)
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# 1. DATA STANDARDIZATION • Read CSV with appropriate headers • Handle duplicate columns (DeKosky) • Map to standard column names: - sequence\_id\_heavy/light - sequence\_heavy/light

#### 2. FASTA GENERATION

- Create BioPython SeqRecord objects
- Write heavy chains → data/fasta-heavy/
- Write light chains → data/fasta-light/

### 3. AIRR ANNOTATION (SADIE)

- Run IgBLAST via SADIE Airr API
- Process heavy chain FASTA files
- Process light chain FASTA files
- Generate AIRR-compliant annotations

# 4. DATA MERGING

- Match heavy/light chains by sequence ID
- Merge annotations with \_heavy/\_light suffixes
- Add metadata from manifest:
  - Run ID, Species, Author
  - B-cell source/type
  - Disease status, etc.

## 5. OUTPUT

- Save as Parquet files in data/parquet-paired/
- Each file contains:
  - Original sequences
  - Complete AIRR annotations
  - Experimental metadata
  - Maintained heavy-light pairing

## **Data Sources**

#### 1. OAS Database

- Location: data/OAS\_paired/
- Format: CSV files with paired heavy/light sequences
- Species: Human antibody sequences
- Manifest: data/oas\_manifest.csv contains metadata for each dataset

#### 2. DeKosky Dataset

- Location: data/DeKosky\_paired/
- Format: CSV files with custom column structure
- Cell Type: Naive B-cells from PBMC
- Special Processing: Requires column renaming due to duplicate headers

#### 3. Leuko Dataset

- Location: data/D326651\_Leuko\_human\_naive.csv
- Cell Type: Naive B-cells from PBMC
- Author: Jonathan Hurtado

# **Processing Pipeline**

# **Step 1: Data Loading and Preparation**

- 1. Read CSV files with appropriate headers (some files have JSON headers requiring special handling)
- 2. Standardize column names:
  - Map to sequence\_id\_heavy, sequence\_id\_light, sequence\_heavy, sequence\_light
  - Handle duplicate column names in DeKosky data

#### **Step 2: FASTA Generation**

- 1. Create FASTA files for heavy and light chains separately:
  - Heavy chains saved to data/fasta-heavy/
  - Light chains saved to data/fasta-light/
- 2. Use BioPython to properly format sequences with IDs

#### Step 3: AIRR Annotation with SADIE

- 1. Run IgBLAST via SADIE's Airr API on each FASTA file
- 2. Generate AIRR-compliant annotations including:
  - V(D)J gene assignments
  - CDR3 sequences
  - o Framework regions
  - Junction analysis

## **Step 4: Paired Data Merging**

- 1. Match heavy and light chains using sequence IDs
- 2. Merge annotations with suffixes \_heavy and \_light

#### 3. Add metadata from manifest:

- Run ID
- Species
- B-cell source (PBMC)
- B-cell type (Naive B-cells)
- Author information
- Disease status
- Other experimental metadata

## **Step 5: Output Generation**

- Save as Parquet files in data/parquet-paired/
- 2. File naming: Uses run ID or dataset identifier
- 3. Format: Apache Parquet for efficient storage and querying

# **Output Structure**

#### Each Parquet file contains:

- Sequence data: Original nucleotide sequences for heavy and light chains
- AIRR annotations: Complete IgBLAST results for both chains
- Metadata: Experimental and sample information
- Pairing information: Maintained heavy-light chain relationships

#### **Technical Details**

# **Dependencies**

- pandas
- BioPython (Bio.Seq, Bio.SeqRecord, Bio.SeqIO)
- SADIE (for AIRR annotation via IgBLAST)

#### **Performance**

- Processing time varies by dataset size
- Example: SRR datasets process in ~20-30 seconds each
- DeKosky datasets: ~4.5 minutes for complete processing

## **Error Handling**

- Checks for existing files to avoid overwriting
- Handles mixed data types in columns
- · Manages memory by deleting dataframes after processing

# Usage Example

## To run the pipeline:

- 1. Ensure all dependencies are installed
  - 2. Place raw data in appropriate directories

- 3. Run the notebook cells sequentially
- 4. Output will be generated in <a href="data/parquet-paired/">data/parquet-paired/</a>

#### **Example Processing Flow**

```
# Process a single OAS file
filename = "ERR4082227"
df = pd.read_csv(f"data/OAS_paired/{filename}_paired.csv")

# Standardize columns
df['sequence_id_heavy'] = df['sequence_id_heavy'].astype(str)
df['sequence_id_light'] = df['sequence_id_light'].astype(str)

# Create FASTA and run SADIE
heavy_df = airr_api.run_fasta(f"data/fasta-heavy/{filename}.fasta")
light_df = airr_api.run_fasta(f"data/fasta-light/{filename}.fasta")

# Merge and save
paired_df = pd.merge(heavy_df, light_df, on='tmp_id', suffixes=('_heavy', '_light'))
paired_df.to_parquet(f'data/parquet-paired/{filename}.parquet')
```

## **Output Example**

Each Parquet file contains ~100+ columns including:

```
Heavy Chain Columns:
- sequence_id_heavy
- sequence_heavy
- v_call_heavy
- d_call_heavy
- j_call_heavy
- cdr3_aa_heavy
- junction_heavy
Light Chain Columns:
- sequence_id_light
- sequence_light
- v_call_light
- j_call_light
- cdr3_aa_light
- junction_light
- . . .
Metadata:
- run
species
bsource
btype
```

- author
- disease
- file\_name

# **Manifest Generation**

A tailored manifest (data/oas\_manifest\_human\_paired.csv) is created containing:

- Only human paired sequences
- Unique author entries
- Sorted by run ID
- Ready for downstream analysis