Translational Bioinformatics for Immunogenomics

Table of contents

W	Welcome 5				
In	ntroduction		6		
1	l Human Leukocyte Antigens		7		
	1.1 Background		7		
	1.2 HLA Class I		7		
	1.3 HLA Class II		7		
	1.4 HLA Nomenclature		7		
	1.5 Functional Divergence		7		
	1.6 HLA Imputation Programs		8		
2	2 Killer Cell Immunoglobulin-like Receptors		9		
	2.1 Background		9		
	2.2 KIR Locus		9		
	2.3 KIR Diversity		12		
	2.4 NK Cell Education		12		
	2.5 KIR Nomenclature		13		
	2.5.1 Inhibitory KIR		13		
	2.5.2 Activating KIR		13		
	2.5.3 Broad KIR Haplotypes		13		
	2.6 KIR Ligand Motifs		13		
	2.7 KIR3DL1 and KIR3DS1		14		
	2.8 KIR Allele Imputation Programs		14		
3	B ERAP	1	16		
4	Epistatic Interactions	1	17		
	4.1 KIR-HLA		17		
В	Bonus	1	18		
5	5 Drug Allergy	1	19		
	5.1 Immediate Drug Allergy		19		
	5.1.1 Skin Testing		19		
	5 1 1 1 Skin Prick Testing		10		

		5.1.1.2 Intradermal Testing
	5.2	Delayed Drug Allergy
		5.2.1 Skin Testing
		5.2.1.1 Intradermal Testing
		5.2.1.2 Patch Testing
	5.3	Specific Drugs
		5.3.1 Antibiotics
		5.3.1.1 Cephalosporings
		5.3.1.2 Fluoroquinolones
		5.3.1.3 Penicillins
		5.3.1.4 Sulfa Antibiotics
		5.3.1.5 Vancomycin
		5.3.2 Buproprion
		5.3.3 Iron
		5.3.4 Local Anesthetics
		5.3.5 Radiocontrast
_		
6		otype Imputation 22
	6.1	Michigan Imputation Server
	6.2	TOPMed Imputation Server
	6.3	Reference Panels
	6.4	Genome Assemblies
7	Bioi	nformatic Best Practices 24
•	7.1	Project Organization
	7.2	Version Control with Git
	7.3	File Naming Conventions
	7.4	Application Containers with Docker
8		senting Your Medical Research 27
	8.1	Font
	8.2	Font Size
	8.3	Word Count
	8.4	Timing
	8.5	Figures
	8.6	References
	8.7	Equipment
		8.7.1 Laptop
		8.7.2 Hub
		8.7.3 USB Drive
		8.7.4 Presentation Remote

9	On	Being a	a Physician-Scientist	30
	9.1	Acade	emic Medicine Jobs	3
		9.1.1	AAMC Faculty Salary Report	3
		9.1.2	Tenure-Track Offer Letters	3
	9.2	NIH I	Loan Repayment Program	3
	9.3	Online	e Resources	3
		9.3.1	Edge for Scholars	3
		9.3.2	NCFDD	3
	9.4	Profes	ssional Organizations	3
	9.5	Sugge	sted Readings	3
		9.5.1	Not Discussed	3
		9.5.2	Publishing Your Medical Research	3
Re	eferer	ıces		32

Welcome

A few key web sources include the Immuno Polymorphism Database and Allele Frequency Net Database.

Introduction

1 Human Leukocyte Antigens

1.1 Background

HLA is located on chromosome 6 in the Major Histocompatibility Complex (MHC).

1.2 HLA Class I

HLA class I molecules are expressed by healthy nucleated cells.

1.3 HLA Class II

HLA class II molecules are expressed by professional antigen-presenting cells (APC)-dendritic cells, macrophages, and B cells.

1.4 HLA Nomenclature

1.5 Functional Divergence

Heterozygosity of HLA class I genes is associated with better outcomes after HIV infection. This is thought to be due to a greater repertoire of HIV peptides presented and cytotoxic T cell response. However, looking at HLA class I allotype alone does not take into account differences in actual peptide repertoire. Viard and O'hUigin developed a metric to measure this difference, termed "functional divergence." Functional divergence predicts the peptide repertoire as a continuum. They showed that greater functional divergence was associated with better HIV outcomes. Functional divergence may be relevant to other diseases where HLA heterozygosity confers advantage, such as infection, vaccination, and immunotherapy.

You can download functional divergence estimates for pairwise combinations of HLA-A, HLA-B, and HLA-C alleles from their article's Supplementary Materials. The functional divergence measure ranges from 0 (i.e., smallest functional divergence) to 1 (i.e., greatest functional divergence).

1.6 HLA Imputation Programs

Name	Programming Language	Input Data	Output	Reference
SNP2HLA	Command line interface	PLINK binary format	HLA class I and II alleles	(Jia et al. 2013)
HIBAG	R	Plink binary format	HLA class I and II alleles	(Zheng et al. 2014)

2 Killer Cell Immunoglobulin-like Receptors

2.1 Background

KIR is located on chromosome 19 (19q13.4) in the Leukocyte Receptor Complex (LRC). KIR is expressed on the surface of Natural Killer (NK) cells and some T cells. KIR do not undergo somatic rearrangement—a key difference from T-cell receptors. KIR interacts with HLA class I—their cognate ligand—to recognize and destroy unhealthy tissue cells while preventing the same from occurring to healthy cells. Therefore, NK cells play a role in fighting infections, resisting some cancers, pregnancy, and preventing autoimmunity. For further reading and references, I highly recommend the review article by Pollock, Harrison, and Norman on the immunogenetics and co-evolution of KIR and HLA class I.

2.2 KIR Locus

Adapted from Pollock, Harrison, and Norman. JACI: In Practice. 2022.

Gene

3DL3

2DS2

2DL2/3

2DL5B

2DS3

2DL1

2DL4

3DL1

3DS1

2DL5A

2DS5

2DS4

2DS1

3DL2

Function

Inhibit.

Activ.

Inhibit.

Inhibit.

Activ.

Inhibit.

Activ.

Inhibit.

Activ.

Inhibit.

Activ.

Activ.

Activ.

Inhibit.

Alleles

228

65

98

47

71

173

112

184

39

44

Allotypes

HLA Class I Ligand Motifs

B7H7

A*11 C1

B46:01 B73:01 C1 C2

PVR

?

C2

HLA-G

Bw4+ HLA-A and Bw4+ HLA-B

 $\mbox{Bw4+ HLA-B}$ and $\mbox{HLA-F}$

PVR

C2

A*11 HLA-C

C2

A3 A11

: Adapted from Pollock, Harrison, and Norman. JACI: In Practice. 2022.

2.3 KIR Diversity

KIR diversity is influenced by gene content variation and sequence variation. Distinct DNA sequences of KIR genes are called "alleles." Distinct polypeptide sequences of KIR genes are called "allotypes." Because different DNA sequences of KIR gene can lead to the same polypeptide, there are more alleles than allotypes for a given KIR gene.

KIR Diversity	
Concept	Definition
Gene	Presence/absence, fusion, duplication
Content	
Variation	
Sequence	May alter ligand affinity or specificity, signal transduction ability, or surface
Variation	expression (e.g., promoter activity, translation, intracellular trafficking)
Allele	Distinct DNA sequence
Allotype	Distinct polypeptide sequence

2.4 NK Cell Education

NK Cell Education		
(i.e., Arming,	Corresponding Pairs of KIR and	Cytotoxicity and other
Licensing)	HLA Class I Ligands	Effector Abilities
Strong	Many	More
Weak	Few	Less

2.5 KIR Nomenclature

2.5.1 Inhibitory KIR

The main role of inhibitory KIR is to prevent cytotoxic NK and T cells from killing tissue cells—unless their HLA class I expression is lost or altered by infection or mutagenesis.

2.5.2 Activating KIR

Activating KIR help identify diseased cells for destruction by cytoxic NK and T cells. Binding of foreign peptides by HLA class I molecules retained by infected cells may be most critical for activating KIR.

2.5.3 Broad KIR Haplotypes

Broad KIR Haplotype	KIR Copy Number Variation	KIR Gene Organization	Activating KIR
A	Relatively stable	Generally	Less
		non-variable	
В	Extensive	Highly variable	More

2.6 KIR Ligand Motifs

Table 2.4: Adapted from Pollock, Harrison, and Norman. JACI: In Practice. 2022.

KIR			
Lig- and			
and			
Mo-			HLA-C
tif	HLA-A Allotypes	HLA-B Allotypes	Allotypes
$\overline{A3/A11}$	A*03, A*11		

KIR Lig- and			
Mo- tif	HLA-A Allotypes	HLA-B Allotypes	HLA-C Allotypes
Bw4	A*23, A*24, A*32	B*07:27, B*08:02, B*08:03, (B13), B*15:13, B*15:16, B*15:17, B*15:23, B*15:24, B*15:36, B*15:43, B*15:67, B*27:01, B*27:02, B*27:03, B*27:04, B*27:05, B*27:07, B*37, B*38, B*40:13, B*40:19, B*44, B*47, B*49, B*51, B*52, B*53, B*56:07, B*57, B*58, B*59	
C1	C*01, C*03, C*07, C*08, C*12:02, C*12:03, C*12:06, C*12:08, C*13, C*14, C*16	B*46, B*73	
C2	C*02, C*03:07, C*04, C*05, C*06, C*12:04, C*12:05, C*12:07, C*14:04, C*15, C*16:02, C*17, C*18		

2.7 KIR3DL1 and KIR3DS1

Because of significant non-allelic recombination in the KIR region, the distinction between KIR genes and alleles can be confusing. Specifically, KIR3DL1 and KIR3DS1 are alleles of the same gene. Of the KIR3DS1 allotypes–3DS1013 and 014–are observed with the greatest frequency in any population.

2.8 KIR Allele Imputation Programs

	Progra	mming		
Name	Lan- guage	Input Data	Output	Reference
PONG	R	PLINK bi- nary for- mat	KIR3DL1/S1 alleles (Global Model includes 51 alleles)	Harrison, 2022
KIR*IMP	Online portal	HAPS/S for- mat	SAMPIRE types: 17 loci (presence/absence and copy number) plus 2 extended haplotype classifications (A and B haplotypes)	Vukcevic, 2015

3 ERAP

ERAP is located on chromsome 5.

4 Epistatic Interactions

4.1 KIR-HLA

Epistatic interactions between KIR and HLA are associated with an kylosing spondylitis (Hanson, 2020)

Bonus

5 Drug Allergy

5.1 Immediate Drug Allergy

5.1.1 Skin Testing

5.1.1.1 Skin Prick Testing

5.1.1.2 Intradermal Testing

5.2 Delayed Drug Allergy

5.2.1 Skin Testing

Table 5.1: Utility of patch and intradermal skin testing for delayed drug allergy reaction types

Reaction	Patch Testing	Intradermal Testing
Maculopapular exanthem	Useful if positive	Useful if positive
(MPE)		
Acute generalized	Useful if positive	Useful if positive
exanthematous pustulosis		
(AGEP)		
Stevens-Johnson	Low sensitivity but	Contraindicated due to
Syndrome/Toxic epidermal	potentially useful if	concern for potential
necrolysis (SJS/TEN)	positive	reactivation
Drug reaction with eosinophilia	Useful if positive	Useful if positive
and systemic symptoms		
(DRESS)		
Fixed drug eruption	Useful if applied to the	Not useful
	site of reaction	
Drug-induced organ injury (e.g.,	Not useful	Not useful
kidney, liver)		

! Important

No delayed skin testing method has 100% negative predictive value.

Table 5.2: Shared characteristics of patch and intradermal testing

Characteristic	Characteristic Details			
Timing	Timing Perform at least 6 to 8 weeks after reaction; and 6 months or later after DRESS			
Concomitant medications	Most medications okay to continue, including anti-histamines and beta-blockers. Should be off of steroids for $$ 1 month or prednisone equivalent dose $$ 10 mg/day			

5.2.1.1 Intradermal Testing

Table 5.3: Characteristics of intradermal testing

Characteristic	Details
Testing site	Volar forearm or extensor upper arm
Testing reagents	Must be sterile; often higher concentrations than those used for
	immediate skin testing
Controls	+ None
	- Saline
Test	+ Papule present
interpretation	- Negative

5.2.1.2 Patch Testing

Table 5.4: Characteristics of patch testing

Characteristic	Details
Testing site	Back of upper arm (needs to be hairless)
Testing reagents	1% and $10%$ of reagent grade product; $10%$ and $30%$ of trade product;
	most commonly used vehicle is petrolatum
Controls	+ None
	- Petrolatum
Shelf-life of patch	Most antibiotics at room temperature are stable for 1 to 3 months;
test mixes	check with USP Pharmacopeia for verification
Patches	Finn chambers (can be aluminum or molded plastic)
Tape	Use hypoallergenic paper tape

Characteristic	Details
Reading	At 48 hours (85% of drugs-if will be positive-are positive by this point); 72 hours; 96 hours; and 1 week
Test interpretation	- Negative - Negative - Doubtful reaction + Weak reaction, erythema ++ Strong reaction, erythema, papules, or vesicles +++ Extreme, bullous, ulcerative

5.3 Specific Drugs

5.3.1 Antibiotics

5.3.1.1 Cephalosporings

5.3.1.2 Fluoroquinolones

5.3.1.3 Penicillins

5.3.1.4 Sulfa Antibiotics

5.3.1.5 Vancomycin

5.3.2 Buproprion

5.3.3 Iron

5.3.4 Local Anesthetics

5.3.5 Radiocontrast

6 Genotype Imputation

6.1 Michigan Imputation Server

The Michigan Imputation Server is a free next-generation genotype imputation platform. You can learn more about the Michigan Imputation Server by visiting their Getting Started documentation. The 1000 Genomes Phase 3 (Version 5) Reference Panel is available on the Michigan Imputation Server.

6.2 TOPMed Imputation Server

The TOPMed Imputation Server is another free next-generation genotype imputation platform developed by the University of Michigan and powered by data from the TOPMed Program investigators. You can learn more about the TOPMed Imputation Server by visiting their Getting Started documentation. The TOPMed Version 3 Reference Panel was released in December 2023.

6.3 Reference Panels

Reference Panel	Genome As- sem- bly	No. of Sam- ples	Sites (chr1- 22)	Chr.	Imputation Server
1000 Genomes Phase 3 (Version 5)	GRCh3	7/ L §194	49,143	,60 5 22,	Michigan Imputation Server
TOPMed (Version 3)	GRCh3	8/11 3<u>8</u>35 97	445,60	X	TOPMed Imputation Server

6.4 Genome Assemblies

The Genome Reference Consortium (GRC) is the main source of human genome assembly data. The most recent human genome assembly version is GRCh38, released in 2013. The "h" in "GRCh" stands for "human." The GRC also maintains genome assembly data for rat (r), mouse (m), zebrafish (z), and chicken (g for gallus). Major updates, called "versions", are released every few years. Minor updates are called "patches" and are released more frequently.

GRCh38 is referred to as "hg38" in the University of California Santa Cruz (UCSC) Genome Browser. The "hg" stands for "human genome." Before the GRCh38 genome assembly, the version numbers of the GRC and UCSC Genome Browser genome assemblies did not match. For example, when the GRCh37 genome assembly was released in 2009, the UCSC Genome Browser version was "hg19." Therefore, to minimize confusion, starting with the GRCh38 genome assembly, the UCSC Genome Browser version number was matched as "hg38."

GRC	UCSC	Year	Genome	
Version	Version	Released	Coverage	Alternate Haplotypes
GRCh37	hg19	2009	$\sim 92.5\%$	3 regions with 9 alternate loci
GRCh38	hg38	2013	95%	178 regions with 261 alternate
				loci

6.5

7 Bioinformatic Best Practices

I recommend the tutorial, "A Reproducible Data Analysis Workflow With R Markdown, Git, Make, and Docker" as a starting point for R-based data analyses (Peikert & Brandmaier, 2021).

7.1 Project Organization

Bash Commands to Create Folder Directory Structure for Your R Project

```
cd </path/to/parent/directory>
mkdir <your-r-project-folder>
cd <your-r-project-folder>
touch README.md
mkdir data doc src bin outputs
```

Once you have downloaded your raw data to your data folder, you should make the contents of the data folder read-only (non-editable) with the following command: chmod u-w -R data/

7.2 Version Control with Git

I recommend the Using Git and GitHub with RStudio Cheatsheet for additional helpful commands.

Verify Git Installation and Version

```
which git # request path to your Git executable
git --version # check your Git version
```

Introduce Yourself to Git

```
git config --global user.name "<username>"
git config --global user.email "<email>"
```

Create a New Repository on GitHub

Go to GitHub to create your new repository, then initialize your repository from the command line.

```
cd </path/to/your-r-project-folder>
echo "# your-r-project-folder" >> README.md
git init
git add README.md
git commit -m "first commit"
git branch -M main
git remote add origin https://github.com/<user.name>/<your-repository>.git
git push -u origin main
```

Add, Commit, and Push Files to Remote Repository

```
git add <file-name>
git commit -m "description"
git push
```

7.3 File Naming Conventions

In your README.md, you should define naming conventions for your project files. The main elements for a file naming convention are metadata, separator, and version tracking. I recommend the File Naming Conventions Worksheet (Briney, 2020) to develop your file naming conventions.

Metadata	Separator	Version Tracking
3 to 5 pieces max (e.g. sample ID, date in ISO 8601 format such	Dashes (-), underscore (_), or camel case (i.e., capitalize each word without spaces)	Numeric (e.g., v01) or Status (e.g., raw, processed)
as YYYY-MM- DD)		

i Example

My naming convention for R Markdown analysis files is: "analysis-YYYY-MM-DD-version.Rmd" where version starts with "v01." This is my first analysis file, "analysis-YYYY-MM-DD-v01.Rmd" $^{\prime\prime}$

7.4 Application Containers with Docker

8 Presenting Your Medical Research

8.1 Font

You should use a sans-serif font like Arial to maximize readability. "Serifs" are extending features at the end of letters. Times New Roman is a serif font.

Table 8.1: Sans-serif versus serif fonts¹

Font	Illustration	Examples	
Sans-serif font	Aal	Bb, Calibri, Helvetica	Cc
Serif font	Aa]	Bb(Times New Roman, Georgia,	$C_{\mathbf{C}}$
Serifs (colored in red)	Aa	Garamond	Cc

8.2 Font Size

 $^{^1}$ Font images are recreated by User:Stannered, original by en:User:Chmod007 - en:Image:Serif and sans-serif 01.png, CC BY-SA 3.0, https://commons.wikimedia.org/w/index.php?curid=2058303

Slide Section	Font Size
Title	36 - 44
Text	24-28
(e.g., Bullets, Figures, Tables)	
References	20-24

8.3 Word Count

The fewer words, the better. A rule to follow is the 7×7 rule: no more than 7 lines and no more than 7 words per line.

8.4 Timing

You should estimate approximately 1 minute per slide.

8.5 Figures

I recommend creating your figures as Scalable Vector Graphics (SVG). The main advantages of the SVG format include always maintaining its resolution and smaller file size than pixel-based image formats (e.g., JPEG).

Some tools that you can use to get started creating SVG include Microsoft PowerPoint (subscription), Adobe Illustrator (subscription), draw.io (free), and Inkscape (free). Draw.io is best for diagrams and flowcharts. Inkscape is better for flexible drawings. Both draw.io and Inkscape are integrated with Bioicons, an open-source extension which includes >1700 icons for scientific illustrations.

In Microsoft PowerPoint, you can create an SVG file by selecting all shapes, right-clicking, choosing "Save as Picture", and then picking "SVG" as the "Save as Type."

8.6 References

Cite references at the bottom of your slides as you present information.

i Format

Last Name. Journal Abbreviation. Year.

8.7 Equipment

8.7.1 **Laptop**

Bring your own laptop to presentations in case there isn't a desktop computer for you to use, or it is not functioning, reliable or frustratingly slow.

8.7.2 Hub

What is worse than not being able to connect your laptop to the correct cable? While a good host for a presentation should have a hub (or dongle if that's your preferred terminology), you can come prepared with your own too–particularly important if you have a laptop with only USB-C ports and no HDMI port.

There are lots of options for hubs. If you are looking for a recommendation, I've found that Anker usually has a selection of high-quality and affordable hubs.

8.7.3 USB Drive

Do you want the entire audience to see your most recent emails when you login to download the PowerPoint you emailed yourself? No. Me either. To avoid this, bring your presentation loaded onto a USB drive, which should ideally have both USB-A and USB-C ports. Or, you can also avoid this by using your own laptop—where the presentation should already be downloaded.

8.7.4 Presentation Remote

I don't feel as strongly about bringing your own presentation remote as your own laptop, hub, and USB drive—but I think it is another piece of equipment to consider. This helps keep you from being tethered to and white knuckling the podium during your talk.

One option to consider is the Logitech Spotlight Presentation Remote—which includes features such as magnification, vibration alerts for time management (e.g., 5 minutes left), 3 hours use from 1 minute of charging, and connection by USB receiver or Bluetooth—in addition to slide advancement.

9 On Being a Physician-Scientist

9.1 Academic Medicine Jobs

9.1.1 AAMC Faculty Salary Report

Looking to get an idea of academic faculty salaries? The annual AAMC Faculty Salary Report compiles academic faculty salaries by rank, degree, department/specialty, medical school type, region, and more. This is often available for free through your university library. Get to know your librarian!

9.1.2 Tenure-Track Offer Letters

What goes into a tenure-track offer letter? The Burroughs Wellcome Fund provides a comprehensive list of offer letter components in their article, "Academic Tenure-Track Offer Letters."

9.2 NIH Loan Repayment Program

NIH Loan Repayment Program

9.3 Online Resources

9.3.1 Edge for Scholars

Edge for Scholars

9.3.2 NCFDD

NCFDD

9.4 Professional Organizations

American Physician Scientists Association

9.5 Suggested Readings

9.5.1 Not Discussed

Not Discussed by Michael Stein

9.5.2 Publishing Your Medical Research

Publishing Your Medical Research by Daniel W. Byrne

References

Jia, Xiaoming, Buhm Han, Suna Onengut-Gumuscu, Wei-Min Chen, Patrick J. Concannon, Stephen S. Rich, Soumya Raychaudhuri, and Paul I. W. De Bakker. 2013. "Imputing Amino Acid Polymorphisms in Human Leukocyte Antigens." Edited by Jianming Tang. *PLoS ONE* 8 (6): e64683. https://doi.org/10.1371/journal.pone.0064683.

Zheng, X, J Shen, C Cox, J C Wakefield, M G Ehm, M R Nelson, and B S Weir. 2014. "HIBAG—HLA Genotype Imputation with Attribute Bagging." *The Pharmacogenomics Journal* 14 (2): 192–200. https://doi.org/10.1038/tpj.2013.18.