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1 **Human thymic T cell repertoire is imprinted with strong convergence to shared sequences**

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35 **Abstract**

36

37 A highly diverse repertoire of T cell antigen receptors (TCR) is created in the thymus by
38 recombination of gene segments and the insertion or deletion of nucleotides at the junctions. Using
39 next-generation TCR sequencing we define here the features of recombination and selection in the
40 human TCR α and TCR β locus, and show that a strikingly high proportion of the repertoire is shared
41 by unrelated individuals. The thymic TCR α nucleotide repertoire was more diverse than TCR β ,
42 with 4.1×10^6 vs. 0.81×10^6 unique clonotypes, and contained nonproductive clonotypes at a higher
43 frequency (69.2% vs. 21.2%). The convergence of distinct nucleotide clonotypes to the same amino
44 acid sequences was higher in TCR α than in TCR β repertoire (1.45 vs. 1.06 nucleotide sequences per
45 amino acid sequence in thymus). The gene segment usage was biased, and generally all individuals
46 favored the same genes in both TCR α and TCR β loci. Despite the high diversity, a large fraction of
47 the repertoire was found in more than one donor. The shared fraction was bigger in TCR α than
48 TCR β repertoire, and more common in in-frame sequences than in nonproductive sequences. Thus,
49 both biases in rearrangement and thymic selection are likely to contribute to the generation of
50 shared repertoire in humans.

51

52 **Keywords:** T cell antigen receptor, TCR repertoire, TCR recombination, thymus, next-generation
53 sequencing

54

55 Abbreviations: T cell antigen receptor (TCR), V (variable), D (diversity), J (joining), CDR3
56 (complementarity-determining region 3), Pgen (generative probability)

57

58 1. Introduction

59 T cell antigen receptor (TCR) is a heterodimeric surface protein, consisting in most cells of α and β
60 chains, while a small minority of cells use γ and δ chains. Both chains are encoded by genes
61 assembled from incomplete segments via somatic recombination during development in the thymus.
62 The TCR β locus contains 47 variable (V), 2 diversity (D) and 13 joining (J) gene segments whereas
63 the TCR α locus contains 42 V and 61 J segments but lacks the D segment. Further diversity is
64 achieved at the gene segment junctions where a number of nucleotides may be removed and
65 palindromic P-nucleotides and non-templated N-nucleotides inserted. Thus, most of the variability
66 in the TCR concentrates in the junctional regions, called complementary determining region 3
67 (CDR3), which also form the main site of antigen recognition (Davis and Bjorkman, 1988).

68
69 The recombination process is capable of creating a high level of diversity. Direct sequencing of
70 TCR β repertoire has measured a lower limit of $1\text{-}3 \times 10^6$ clonotypes, whereas a mathematical
71 estimator suggested a total repertoire of about 100×10^6 unique clonotypes (Qi et al., 2014). We have
72 recently measured the lower limit of thymic TCR diversity in pediatric samples to be 10.3×10^6 for
73 TCR β and 3.7×10^6 for TCR α clonotypes, and statistical modelling suggested the total repertoire to
74 consist of $40\text{-}70 \times 10^6$ and $60\text{-}100 \times 10^6$ clonotypes for TCR β and TCR α respectively (Vanhanen et
75 al., 2016). The pairing of TCR α to TCR β has been studied but little. A sequencing of a limited TCR
76 subset showed that on the average each TCR β chain can bind to at least 24 different TCR α chains
77 (Arstila et al., 1999), while a recent large scale single-cell analysis suggested that the pairing is
78 more limited than would be compatible with a fully stochastic process (Grigaityte et al., 2017). The
79 full TCR $\alpha\beta$ repertoire thus consists of at least tens of millions of different receptors.

80
81 To date the human thymic TCR repertoire has been studied very little. The TCR β locus is
82 rearranged first and is subject to relatively stringent allelic exclusion. However, TCR β locus may

83 also be rearranged in cells destined to the $\gamma\delta$ T cell lineage, which may account for a part of the
84 nonfunctional TCR β repertoire. Since most recombination events will result in an out-of-frame
85 sequence, the functionality of the rearranged TCR β chain is ensured by pairing with a surrogate
86 TCR α chain, the preT α . The cells capable of signaling through pre-TCR then proliferate before
87 recombination begins in the TCR α locus (von Boehmer et al., 1998). Unlike the TCR β locus, in
88 TCR α recombination both alleles are rearranged simultaneously, until a functional TCR $\alpha\beta$ is
89 expressed, stopping the recombination. Thus, in a large proportion of cells both TCR α loci are
90 rearranged, although only one is likely to produce a functional protein chain (Casanova et al.,
91 1991). The newly generated TCR $\alpha\beta^+$ cells are then subjected to positive and negative selections,
92 which remove cells incapable of interacting with HLA molecules or displaying too strong affinity to
93 self-antigens (Klein et al., 2014). Overall, only an estimated 3-5% of the developing thymocytes
94 survive the selection process to form the mature peripheral repertoire (Egerton et al., 1990; Yates,
95 2014).

96

97 In the present study, we characterize the composition of the thymic TCR β and TCR α repertoire,
98 identifying differences in the two chains related to their biology. Our data also show a strikingly
99 strong convergence to shared repertoire in unrelated individuals.

100 **2. Materials and Methods**

101 The study was approved by the Pediatric Ethical Committee of the Helsinki University Hospital and
102 parents gave a written informed consent. Thymus samples were obtained from eight
103 immunologically healthy children undergoing a corrective operation for congenital cardiac defects
104 (donors A-D and donors 1-4). Additionally, a peripheral blood sample was drawn from donors 1-4
105 during the operation. The donors were 7–244 days old and 2/8 were female (Table 1). Two of the
106 subjects (donors A and B) were monozygotic twins. The impact of genetics on the repertoire has
107 been analyzed in detail elsewhere (Heikkila et al., 2020). All thymus samples appeared
108 macroscopically normal. Thymocyte populations from donors B-D were analyzed with flow
109 cytometry for expression of CD4, CD8, TCR $\alpha\beta$, TCR $\gamma\delta$, CD3 and CD69.

110
111 From each subject, an aliquot of 10–30 million thymocytes and from donors 1-4 an aliquot of 0.5
112 mL peripheral blood was used for sequencing both TCRAD and TCRB repertoire. Thymocytes
113 were extracted mechanically from the tissue. To remove red blood cells blood samples were treated
114 with Gibco™ACK Lysing Buffer (Thermo Fisher Scientific, Massachusetts, USA), according to the
115 manufacturer’s orders. Genomic DNA was extracted from pelleted cell samples with QIASymphony
116 (Qiagen, Germany) according to the manufacturer’s instructions. The TCR α and TCR β CDR3
117 regions were amplified and sequenced from a standardized quantity of quality-controlled DNA
118 using ImmunoSEQ assay (Adaptive Biotechnologies, Seattle, USA). In summary, the sequencing
119 assay consists of a multiplex PCR system to amplify the rearranged CDR3 regions from the DNA
120 samples at a length that is sufficient to subsequently identify the VDJ and VJ regions spanning each
121 unique CDR3 α and CDR3 β regions, respectively. Amplicon sequencing was performed with
122 Illumina platform. TCR α and TCR β gene segment definitions were obtained from IMGT database
123 (www.imgt.org). Primer bias was corrected as previously described (Vanhanen et al., 2016) and the
124 resulting data filtered and clustered using both the relative frequency ratio between similar clones

125 and a modified nearest-neighbor algorithm to remove both PCR and sequencing errors. All
126 sequences are available at immuneACCESS database provided by Adaptive Biotechnologies
127 (clients.adaptivebiotech.com/immuneaccess).

128
129 The TCR sequence analysis was performed using the immunoSEQ ANALYZER 3.0 (Adaptive
130 Biotechnologies, Seattle, USA), VDJTools software (Shugay et al., 2015) and in-house scripts for
131 computing languages R (www.r-project.org) and python 2.7 (www.python.org). The in-house
132 scripts generated for this study are published in Supplements 1&2. The similarity of two sets of
133 unique or total sequences was assessed calculating the Jaccard index, which is defined as the size of
134 the intersection of two data sets (A and B) divided by the size of their union: $J(A, B) = \frac{|A \cap B|}{|A \cup B|}$. The
135 abundance based Jaccard index was defined as $J_{abund} = UV/(U+V-UV)$, where U is the total relative
136 abundance of shared sequences in sample A and V the total relative abundance of shared sequences
137 in sample B (Chao et al., 2006). The CDR3 nucleotide sequences were extracted separately for in-
138 frame and nonproductive sequences and subsequently the generative probabilities were calculated
139 using the OLGA software (Sethna et al., 2019).

140

141 **3. Results**

142 **3.1. TCR α and TCR β repertoires differ in diversity and productivity**

143 Thymus samples were collected from eight pediatric patients (donors A-D and donors 1-4), two of
144 whom were monozygotic twins (donors A and B; Table 1). Flow cytometric analysis was performed
145 for donors B-D and showed a normal distribution of CD4 and CD8 double-negative (DN), double-
146 positive (DP), and single-positive (SP) thymocytes as well as normal pattern of TCR $\alpha\beta$ and TCR $\gamma\delta$
147 expression (Figure 1A). Postselection thymocytes were defined as DP $CD3^{high}CD69^{+}$, CD4SP or
148 CD8SP (Swat et al., 1993; Yamashita et al., 1993). On the average, $23.1\pm 3.7\%$ of total thymocytes
149 represented postselection and $76.9\pm 3.7\%$ preselection population (Figure 1B).

150

151 Sequencing of thymic TCRs yielded 1.2×10^5 - 1.6×10^6 (mean 810 000) unique TCR β clonotypes of
152 which $78.8\pm 2.7\%$ were in-frame, $19.3\pm 2.4\%$ were out-of-frame and $2.1\pm 0.5\%$ contained a
153 premature stop-codon (Fig. 1C). Consistent with our previous estimation on thymic TCR diversity,
154 the TCR α diversity was higher than TCR β diversity, with 1.3 - 7.6×10^6 (mean 4.1×10^6) unique
155 clonotypes per sample (Vanhanen et al., 2016). However, the productivity in TCR α was much
156 lower, as only $30.8\pm 0.8\%$ of the unique clonotypes were in-frame. Of the unique TCR α clonotypes
157 $66.0\pm 0.6\%$ were out-of-frame and $7.0\pm 4.5\%$ contained a premature stop-codon (Fig. 1C). As the
158 sequencing assay is based on genomic DNA, it also provides a quantitative estimate of the number
159 of total genomes with rearranged TCR segments in the sample.

160

161 A small blood sample from donors 1-4 was sequenced simultaneously with the thymus samples,
162 producing an average of 84 000 unique TCR β clonotypes and 150 000 unique TCR α clonotypes. In
163 the TCR β repertoire, the fractions of in-frame and nonproductive clonotypes remained essentially
164 similar to that in the thymus (Figure 1D). In the TCR α repertoire, the fraction of in-frame

165 clonotypes was higher in the blood samples than in the thymus ($38.5\pm 1.4\%$ vs. $30.8\pm 0.8\%$; Figure
166 1D).

167

168 To estimate the convergence of distinct nucleotide clonotypes to identical amino acid chains we
169 calculated the nucleotide-to-amino acid-ratio for each sample. The majority of amino acid chains in
170 the TCR β repertoire were encoded by a single nucleotide clonotype, the nucleotide-to-amino acid-
171 ratio being for unique in-frame clonotypes 1.06 ± 0.03 in the thymus and 1.05 ± 0.02 in the periphery.
172 In the TCR α repertoire the number of unique nucleotide clonotypes converging to the same amino
173 acid chain was higher than in the TCR β repertoire, particularly in the thymus (ratio 1.45 ± 0.13) but
174 also to some degree in the periphery (ratio 1.18 ± 0.01).

175

176 **3.2. The V and J segment usage is biased before thymic selections**

177 Previous studies of peripheral repertoire have shown a biased usage of V and J genes in healthy
178 subjects. Similarly, in thymus the use of V gene elements was uneven, and the same segments were
179 favored in each individual both in thymus and in blood (e.g. TRBV5-1, TRBV27-01 and TRAV21-
180 1, TRAV29-1; Supplement 3). Similar findings were also obtained for J gene usage (Supplement 4).
181 The biased V and J gene usage pattern was largely observed both in the in-frame and nonproductive
182 repertoire, indicating that it is due the recombination process rather than selection (Figure 2).
183 Consistent with our previous study (Heikkila et al., 2020), the samples from the monozygotic twins
184 A and B clustered together, indicating a genetic component in V and J gene usage. Interestingly, in
185 the TCR α repertoire, the gene segment usage clustered thymic and peripheral blood samples mainly
186 according to the sample type and not the identity of the donor (Figure 2A). In the TCR β repertoire,
187 in contrast, the gene segment usage clustered together blood and thymus samples taken from the
188 same donor (Figure 2B).

189

190 Some of the gene segment bias might be caused by thymic generation of semi-invariant T cell
191 subsets, such as natural killer T cells (NKTs) or mucosal-associated T cells (MAITs). Human NKTs
192 prefer TRAV10/TRAJ18 combination and MAITs use invariable TRAV01-02/TRAJ33-01
193 combination. The β chain usage is less restricted, but with a preference of TRBV25 for NKTs and
194 TRBV6 and TRBV20 for MAITs. In our data none of the semi-invariant α chains was dominant
195 whereas some MAIT-associated TRBV6 genes were found at an elevated frequency. However,
196 these TRBV segments are also ubiquitously used by conventional variable T cells (Tickotsky et al.,
197 2017).

198
199 TCR δ gene segments are embedded within the TCR α locus and $\alpha\beta$ and $\gamma\delta$ lymphocytes may use
200 both TCR α and TCR δ gene segments in an overlapping manner (Verschuren et al., 1998). Since the
201 thymocytes we analyzed were not sorted, and the sequencing protocol included primers specific for
202 the entire TCRAD locus, we obtained a mixture of TCR α and TCR δ sequences. In the thymus, the
203 frequency of $\gamma\delta$ TCR $^+$ thymocytes, as measured by flow cytometry, was $0.80\pm 0.20\%$. However, the
204 frequency of unique clonotypes using a combination of TRDV and TRDJ was $1.1\pm 0.18\%$. In the
205 peripheral blood the frequency of TRDV-TRDJ combinations was slightly higher ($1.7\pm 0.96\%$ of
206 the unique clonotypes). We also identified relatively frequent combinations of TRDV to TRAJ
207 ($2.4\pm 0.20\%$ of the unique clonotypes), whereas sequences using a combination of TRAV and
208 TRDJ were rare both in thymus and in periphery (Table 2).

209

210 **3.3. CDR3 region length reflects recombination and selection events**

211 The TCR β chain comprises V, D, and J segments, whereas the TCR α chain lacks D segments and
212 thus contains only one junctional site. This difference was reflected in the higher number of non-
213 templated nucleotide insertions in the TCR β than in the TCR α sequences with an average of 9.3 vs.
214 3.9 nucleotides in thymic and 7.0 vs. 3.7 in peripheral in-frame repertoires (Figure 3A). The

215 nonproductive sequences cannot be subject to TCR-mediated selection, and thus represent the non-
216 selected product of the recombination process. Consistent with the previously reported shortening
217 of CDR3 during thymic selection (Matsutani et al., 2011; Niemi et al., 2015; Yassai and Gorski,
218 2000), the mean CDR3 length was shorter in the in-frame rearrangements (41.6 base pairs (bp) for
219 TCR α and 45.7 bp for TCR β) than in the nonproductive rearrangements (41.9 bp for TCR α and
220 46.3 bp for TCR β) in the thymus (Figure 3B). In the peripheral in-frame repertoire, the CDR3
221 regions were still shorter (41.4 bp for TCR α and 43.4 bp for TCR β).

222

223 **3.4. Pgen distributions differ in TCR α and TCR β repertoires**

224 In the process of V(D)J gene segment recombination and insertion of random nucleotides between
225 gene segments some sequences are generated more readily while the generation of others is more
226 unlikely. We used OLGA software to calculate the generative probabilities (Pgen) in the TCR α and
227 TCR β repertoires (Sethna et al., 2019). For a large majority of nonproductive sequences we
228 obtained Pgen values 0, probably because the OLGA calculations are based on amino acid rather
229 than nucleotide sequences and CDR3 amino acid definition remains ambivalent for nonproductive
230 sequences. For the thymic in-frame sequences the Pgen was higher for TCR α (average Pgen 1.57e-
231 7) than for TCR β (average Pgen 1.34e-9) repertoire, a finding likely due to the lower junctional
232 complexity in TCR α chains. The same was observed in the peripheral repertoires (average Pgen for
233 TCR α 1.56e-7 and for TCR β 3.62e-9). In the TCR α repertoire, the thymic and peripheral Pgen
234 averages and distributions were largely identical, while for the TCR β the thymic repertoires had
235 lower Pgen values than the peripheral repertoires (1.34e-9 vs. 3.62e-9; Figure 4).

236

237 **3.5. Overlap of thymic clonotypes between two individuals**

238 Despite the high diversity of the junctional CDR3 sequences, a considerable overlap of peripheral
239 TCR repertoires between different individuals has been reported (Shugay et al., 2013). In our

240 thymic samples, a substantial fraction of TCR sequences were shared between two individuals, and
241 some of the TCR α and TCR β clonotypes were shared even between multiple individuals (Figure
242 5A). This phenomenon was more marked in the TCR α than TCR β repertoire. Indeed, in the samples
243 1-4, in which the sequencing depth was shallower, there were no TCR β clonotypes shared by all
244 four donors.

245

246 To estimate the fraction of thymic repertoire shared by two individuals, we used the Jaccard index
247 (JI), calculated as the intersection of two samples divided by the union of the samples, with a
248 maximum index of 1 for fully overlapping repertoires. In the nonproductive TCR β clonotypes the JI
249 was low (mean JI $6.3e-5$), but increased clearly in the in-frame repertoire (mean JI $4.6e-4$). When
250 unique amino acid CDR3 regions were analyzed, the shared fraction was higher still (mean JI
251 0.013; Figure 5B). In the TCR α repertoire, the shared fraction was generally higher than in the
252 TCR β repertoire, and in the nonproductive clonotypes the mean JI was 0.029. A small but
253 consistent increase to mean JI of 0.032 was found in the in-frame repertoire. In the unique amino
254 acid CDR3 regions the shared fraction was again clearly higher (mean JI 0.10; Figure 5B). As
255 previously reported (Heikkila et al. 2020), comparison of the twins A and B produced slightly
256 higher JIs than the other pairs. In general, samples 1-4 were sequenced to a lesser depth than
257 samples A-D, affecting the observed number of shared clonotypes, and the JI values were
258 consequently smaller. However, the increasing trend in JI from nonproductive to in-frame and
259 amino acid sequences was clear in all samples.

260

261 The shared sequences contained fewer non-templated insertions than the individual private
262 repertoires. The average number of non-templated insertions in was 1.4 and 2.6 respectively for
263 shared in-frame and nonproductive TCR β clonotypes. In TCR α the shared in-frame clonotypes
264 contained on the average 1.4 and nonproductive 1.6 insertions. Also, the Pgen was higher in the

265 shared repertoire compared to the full repertoires. In the in-frame repertoire the average Pgen for
266 unique shared in-frame TCR β clonotypes was 4.71e-8 and in the full repertoire 1.34e-9. For in-
267 frame TCR α clonotypes the difference in Pgen between shared (2.38e-7) and full (1.57e-7)
268 repertoires was smaller than for TCR β but still distinct.

269

270 Since our sequencing method uses genomic DNA instead of messenger-RNA as starting material, it
271 has been optimized for quantitative analysis and provides us with a reasonable estimate of the
272 clonal abundance (Robins et al., 2009; Vanhanen et al., 2016). Thus, the analysis of the shared
273 fraction of total genomes reflects the actual size of repertoire common to different individuals. For
274 total genomes, a similar increasing trend in JIs from nonproductive to in-frame and to amino acid
275 repertoires was observed as seen for unique sequences. In total in-frame nucleotide genomes the
276 mean JI for TCR α repertoire was 0.083 and for TCR β repertoire 0.00063. In total amino acids the
277 shared part of the repertoire was extremely large (mean JI 0.30 for TCR α and 0.026 for TCR β ;
278 Figure 5C). In percentages, on the average, of the total TCR β amino acid repertoire of any given
279 individual 6.1% was also found in the repertoire of another individual (range 1.55-11.4%). In the
280 TCR α repertoire the overlap in percentages was strikingly high (mean 46.7%, range 32.6-62.7%;
281 Supplement 5).

282

283 **3.6. Sharing of high abundance clones**

284 To analyze the relationship between clone size and the likelihood of sharing, we calculated the
285 Jaccard indexes for the most abundant 1%, 2%, 5%, 10%, 20% and 50% of clones. For this analysis
286 samples 1-4 were excluded, because the relatively shallow sequencing produced very little overlap
287 among the top 1-5% clonotypes. In TCR α repertoire we observed a clear correlation between the
288 sharing and the clonotype abundance. JI values were clearly highest in the top 1-2% most abundant
289 clonotypes and decreased gradually when less abundant clonotypes were included (Figure 6A). In

290 contrast, there was no similar correlation in the TCR β repertoire and the interindividual variation in
291 JIs among the top 1% most abundant clones was very wide (Figure 6A). The number of non-
292 templated nucleotide inserts also showed a correlation with the sharing among highly abundant
293 clones. Non-templated inserts were rare among the most abundant shared clones. In the TCR α
294 repertoire the average number of inserts in the shared repertoire increased steadily with the analysis
295 of less abundant clonotypes (Figure 6B). In the TCR β repertoire the number of inserts was typically
296 zero among the top 2% most abundant shared repertoire and increased abruptly for the top 5-50%
297 most abundant clonotypes (Figure 6B).

298

299 **3.7. Sequence overlap in the peripheral samples**

300 Despite the clearly smaller number of cells analyzed, clonotype sharing was also observed in the
301 peripheral blood. Similarly to the thymus, sharing was higher in the TCR α than in the TCR β
302 repertoire and some clonotypes were shared between all four samples (Figure 7A). Also in the
303 peripheral samples, sharing was lowest in the nonproductive nucleotide repertoire, increased in the
304 in-frame nucleotide and even more so in the amino acid CDR3 repertoires (Figure 7B-C).

305 4. Discussion

306 Until recently, our understanding of the human thymus has been largely based on extrapolation
307 from circulating repertoire and from murine studies. However, studies on organ donors combined
308 with high-throughput techniques and next-generation sequencing have begun to provide
309 information on the various types of cells in the human thymus (Park et al., 2020; Thome et al.,
310 2016). A single-cell sequencing study coupled with TCR $\alpha\beta$ profiling identified approximately 200
311 000 individual lymphoid cells among 24 fetal and mature thymi and showed a biased V(D)J usage
312 originating from recombination and modified by selection (Park et al., 2020). We have previously
313 estimated the total thymic TCR diversity to be 60-100x10⁶ for TCR α and 40-70x10⁶ for TCR β
314 repertoire and thus currently beyond the coverage of single-cell experiments (Vanhanen et al.,
315 2016). Our current data from eight pediatric thymi comprises a total of 161 million TCR α reads and
316 55 million TCR β reads, representing the most extensive characterization of the thymic TCR
317 repertoire so far. Although our analysis was performed on unsorted cells and thus allows little
318 conclusions on the developmental stage and functionality of the TCRs, the large scale provides an
319 opportunity to compare specific features of TCR α and TCR β repertoires and, particularly, to
320 measure thymic repertoire overlap across individuals.

321

322 As previously reported for peripheral blood samples and recently for thymus as well (Park et al.,
323 2020; Quiros Roldan et al., 1995; Zvyagin et al., 2014) the usage of V and J gene segments is
324 clearly biased in the thymus. The same gene segments were dominant in every individual, in both
325 the TCR α and TCR β chains. Some of this bias has been ascribed to selection by HLA molecules,
326 which interact with protein loops encoded by the germ-line parts of TCR V genes (Huseby et al.,
327 2005; Rudolph et al., 2006; Wu et al., 2002). However, the same biased usage was also observed in
328 the nonproductive repertoire, which cannot be subjected to selection by antigen-HLA complexes.
329 This suggests that the bias is partly generated in the recombination itself. We have previously

330 reported that genetic factors influence the gene segment usage in the thymus, a finding confirmed
331 here with an increased number of samples. Our data also show that the use of TCRD elements in $\alpha\beta$
332 T cells is common, with ca. 6% of thymic sequences containing TCRD gene segments, while the
333 frequency of $\gamma\delta$ TCR+ thymocytes was less than 1%. However, combining TCRAV to TCRDJ
334 seems to be largely prevented.

335

336 Despite the structural and functional similarity of the two TCR chains, the generation of TCR α and
337 TCR β repertoire has several differences, which are also reflected in our data. First, the number of
338 non-templated nucleotide insertions was much higher in the TCR β locus, most likely explainable by
339 the fact that, unlike TCR α chain, TCR β chain undergoes two recombination events (D to J followed
340 by V to DJ). This is also displayed by the slightly longer CDR3 region length and lower calculated
341 Pgen in TCR β than in TCR α repertoire. Second, the number of non-templated inserts and CDR3
342 length were lower and respectively Pgen was higher in the peripheral than in thymic samples in
343 TCR β repertoire while in TCR α these features remained relatively similar in thymus and periphery.
344 Third, the fraction of in-frame rearrangements was higher in TCR β than in TCR α locus (78.8% vs.
345 30.8% in the thymus). This reflects the difference in allelic exclusion in TCR β and TCR α locus. In
346 TCR β locus the exclusion is strict, whereas both TCR α loci are rearranged simultaneously and a
347 large fraction of cells will end up with a nonfunctional rearrangement in the other TCR α locus
348 (Borgulya et al., 1992; Casanova et al., 1991). The frequency of nonproductive sequences in our
349 samples is also increased by the presence of immature thymocytes not yet subjected to TCR-
350 mediated selection, and the ongoing TCR α locus rearrangement in some of the cells. Furthermore,
351 the repertoire overlap was much higher in TCR α than TCR β repertoire, consistent with previous
352 analyses (Khosravi-Maharlooei et al., 2019; Zvyagin et al., 2014), but here shown in a large-scale
353 analysis of thymic repertoire. Although we obtained fewer TCR β than TCR α sequences, it is clear

354 that the higher sequence overlap in TCR α compared with TCR β is mostly biological and not due to
355 differences in sequencing depth, a finding also confirmed by others.

356

357 Indeed, the remarkably high degree of clonal sharing between individuals is the most interesting
358 observation in our current data. Here, it must be noted that two of our donors were monozygotic
359 twins, which introduces a bias to the analysis. However, these two samples only shared a
360 marginally higher fraction of sequences than the unrelated samples (Heikkila et al., 2020). This is
361 largely consistent with a recent analysis of the peripheral repertoire in three pairs of identical twins,
362 which concluded that there was no difference between the twins and unrelated donors in the sharing
363 of CDR3 sequences (Zvyagin et al., 2014).

364

365 Given the enormous diversity of possible TCRs, the expected likelihood to detect identical
366 receptors in two individuals is practically nonexistent. Still, previous studies of inbred mouse lines
367 have reported that roughly 30% of the peripheral TCR β repertoire is shared (Bousso et al., 1998;
368 Furmanski et al., 2008). More recent studies have used next-generation sequencing methods,
369 analyzing much larger numbers of sequences. Sequencing of TRBV12-4/TRBJ1-2 expressing
370 peripheral blood CD8⁺ T cells in four unrelated healthy donors yielded in average 29 000 unique
371 clonotypes per individual and the overlap of unique amino acid CDR3 sequences was 3.8–9.8%
372 (Venturi et al., 2011). Zvyagin et al. measured the overlap of both TCR β and TCR α repertoires in
373 three pairs of monozygotic twins reaching an overlap of 3–10% and 10–26.5% of unique amino
374 acid CDR3 clonotypes in TCR β and TCR α repertoires, respectively, without higher similarity
375 between the twins than unrelated pairs (Zvyagin et al., 2014). It was also estimated that if the
376 predicted peripheral TCR β diversity of 5×10^6 unique sequences was entirely sequenced, the CDR3
377 overlap between two individuals would reach 44.1% in the amino acid and 3.6% in the nucleotide
378 repertoire (Shugay et al., 2013).

379

380 In our thymus data, taking into account the clonal abundances of the clonotypes, the average
381 fraction of sequences found in any other donor for the total TCR β repertoire was 0.2% for in-frame
382 nucleotide chains and 6.1% for amino acid CDR3 chains. In the TCR α repertoire, the sharing was
383 much higher: an average of 15.7% total in-frame nucleotide and 46.7% total amino acid CDR3
384 chains were shared between any two donors. Furthermore, in the TCR α repertoire the overlap
385 showed a strong correlation with clone abundance, whereas the same was not true of the TCR β
386 repertoire. The average number of non-templated inserts was also lower among the most abundant
387 clonotypes both for TCR α and TCR β repertoire. Together, these data suggest that some TCR α
388 sequences are generated easily and preferred across different unrelated individuals.

389

390 Notably, the surprisingly high repertoire overlap in thymus was directly measured from samples of
391 10 million thymocytes, taken from an organ with an estimated 50 billion cells (Ganusov and De
392 Boer, 2007; Rodewald, 2008). Recent analyses have shown that the degree of sharing is correlated
393 with the size of the sample sequenced (Campregher et al., 2010; Putintseva et al., 2013; Shugay et
394 al., 2013; Venturi et al., 2011). It is thus possible that exhaustive sequencing of the thymic
395 repertoire would reveal an even higher proportion of shared sequences. Indeed, it may be that TCR
396 repertoire is really individualized only by α -to- β pairing, although even this may be less stochastic
397 than previously assumed (Grigaityte et al., 2017).

398

399 It is clear that some of this sharing reflects convergent recombination, i.e., the recombination
400 machinery favoring certain gene segments and particular types of CDR3 sequences. Previous
401 analysis of peripheral repertoire has shown that shared sequences have relatively few nucleotide
402 additions and are generally closer to germline sequences (Pogorelyy et al., 2017; Quigley et al.,
403 2010; Venturi et al., 2008a; Venturi et al., 2006). This is also seen in our thymus samples, where the

404 shared sequences had on average fewer nucleotide insertions and higher calculated Pgen than the
405 repertoire in general. This implies that some junctional sequences are easier to generate and
406 therefore appear repeatedly, and their high frequency may therefore not require peripheral
407 expansion (Venturi et al., 2008b).

408

409 However, our quantitation showed a strong enrichment of the shared repertoire the further the
410 sequences receded from the recombination process. In every donor pair the shared fraction was
411 higher in the in-frame than in the nonproductive repertoire and higher still in amino acid sequences
412 and total number of genomes. This was particularly striking in the TCR β repertoire, in which the
413 average JI increased from 6.3×10^{-5} in the nonproductive repertoire to 0.026 in total amino acid
414 genomes, or by a factor of ~ 400 . In the TCR α chain the increase was by a factor of ~ 10 , from 0.026
415 to 0.30. Since the nonproductive nucleotide sequences are not subject to any form of TCR-mediated
416 selection, this enrichment indicates that a substantial fraction of the clonal sharing is due to antigen-
417 driven selection in the thymus.

418

419 In the periphery, although shared clones specific to defined antigens have been described, the
420 antigen-dependent selection seems in general to lead to divergence in the repertoire. Analysis of
421 naive and memory CD8 $^{+}$ T cells found fewer shared clones in the latter, antigen-experienced
422 population, while a comparison of preterm neonates with adults showed that the shared fraction of
423 TCR β (CDR3 amino acid chains) decreased from 8% to 1% (Carey et al., 2017). Similarly, donors
424 in younger age groups shared a larger fraction of TCR β repertoire than older individuals and while
425 TCR β repertoires in young are similarly high in diversity, with age clonal expansions accumulate
426 and the individual repertoires develop to divergent directions (Britanova et al., 2014; Britanova et
427 al., 2016). In our data the shared fraction of CDR3 amino acid sequences in the peripheral blood

428 was 5.3% in the TCR β and 17.1% in the TCR α repertoire, in donors ranging from 7 days to 5
429 months of age.

430

431 A further point relates to the transitory nature of thymic function. Since thymus is a primary
432 lymphoid organ constantly producing new T cells, any given clone will spend only a limited time in
433 the thymus before either failing selection and dying or maturing and emigrating to periphery. The
434 repertoire might thus also be expected to be transitory, with a different snapshot of the repertoire
435 obtained at different points in time. In contrast, the high degree of interindividual clonal sharing
436 suggests by extension that at different time points a given thymus is producing similar clones.
437 Indirectly, our results imply that although the thymic T cell population and TCR repertoire is
438 transitory, the clonal composition of human thymus is surprisingly stable.

439

440 In conclusion, our study provides the first detailed characterization of the human thymic TCR α and
441 TCR β repertoire, showing similarities and differences in the features of these two TCR chains. We
442 also show an unexpectedly high overlap of thymic TCR repertoire between unrelated donors,
443 especially in the TCR α chain. Moreover, our data indicate that this convergence is substantially
444 driven by thymic selection. Finally, it must be noted that the specificity of any TCR is determined
445 by α -to- β pairing, which our data do not address. As shown by Grigaityte et al., novel technology is
446 finally allowing this part of the repertoire to be analyzed, as well (Grigaityte et al., 2017).

447

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455

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457

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610

611

612 **Tables**

613 **Table 1.** Description of the samples. The details of each sequenced sample and the numbers of
614 obtained unique clonotypes and total reads per sample for TCR α and TCR β repertoires.

Sample	Age (days)	Sex	TCR α		TCR β	
			Unique	Total	Unique	Total
Thymus A	243	M	6 907 422	39 865 283	1 254 760	8 431 833
Thymus B	244	M	7 578 104	45 335 572	1 540 161	11 558 445
Thymus C	225	F	5 347 824	30 309 225	1 568 528	23 581 729
Thymus D	126	M	6 743 495	36 762 724	1 462 150	11 159 872
Thymus 1	7	M	2 089 557	3 179 774	223 725	237 063
Thymus 2	52	M	1 262 845	1 747 487	173 368	182 356
Thymus 3	107	M	1 289 728	2 158 043	138 544	142 903
Thymus 4	156	F	1 419 013	1 848 851	122 195	128 228
Average			4 079 749	20 150 870	810 429	6 927 804
Blood 1	7	M	138 159	154 682	77 868	82 418
Blood 2	52	M	109 171	123 523	69 875	73 945
Blood 3	107	M	180 100	245 126	104 236	134 110
Blood 4	156	F	167 266	199 326	82 550	88 901
Average			148 674	180 664	83 632	94 844

615

616

617 **Table 2.** Mean frequency (%) of V δ and J δ segments in the TCR α repertoire

		Thymus	Peripheral blood
Vδ-Jδ	Unique	1.09	1.71
	Total	2.06	3.83
Vδ-Jα	Unique	2.39	2.71
	Total	3.81	3.84
Vα-Jδ	Unique	0.36	0.42
	Total	0.41	0.51

618

619

620 **Figure captions**

621

622 **Figure 1.** Analysis of the thymocyte subsets and repertoire productivity. The fraction of TCR $\alpha\beta$ +
623 and TCR $\gamma\delta$ + in thymocytes and the distribution of CD4 and CD8 among TCR $\alpha\beta$ + thymocytes in a
624 representative thymus sample (donor C) with the applied backgating (A). The distribution of CD4
625 and CD8 expression in thymocytes and the fraction of CD3^{high}CD69+ cells in CD4+CD8+ double
626 positive thymocytes (donor C) with the applied backgating (B). The fraction of sequences in-frame,
627 out-of-frame, or containing a premature stop codon among unique TCR α and TCR β clonotypes for
628 thymic (C) and peripheral TCR repertoires (D).

629

630 **Figure 2.** The V gene usage in in-frame and nonproductive repertoires. The heatmaps display the
631 frequencies of different V gene segments and the attached dendrograms show the clustering of the
632 samples in in-frame and nonproductive TCR α (A) and TCR β repertoires (B).

633

634 **Figure 3.** The number of non-templated insertions and the CDR3 lengths. The graphs show
635 the average and 95% confidence interval of the number of non-templated nucleotide insertions (A)
636 and of CDR3 lengths (B) in thymic and peripheral blood TCR α and TCR β repertoires for in-frame
637 and nonproductive sequences.

638

639 **Figure 4.** The generation probability (Pgen) calculated with OLGA software. Thymic and
640 peripheral Pgen distribution plotted against probability density in the in-frame TCR α and TCR β
641 repertoires for a representative thymus-blood pair (donor 1).

642

643 **Figure 5.** Sequence overlap between thymus samples. Venn diagrams show the overlap of unique
644 in-frame clonotypes separately for thymus samples A-D and 1-4 (A). Individual Jaccard indexes

645 (JI) between each thymus sample for nonproductive, in-frame and amino acid repertoires among
646 unique clonotypes (B) and total genomes (C). Monozygotic twins A and B are identified as open
647 circles, filled circles represent the JI between unrelated individuals. The average JI and the 95%
648 confidence interval are shown.

649

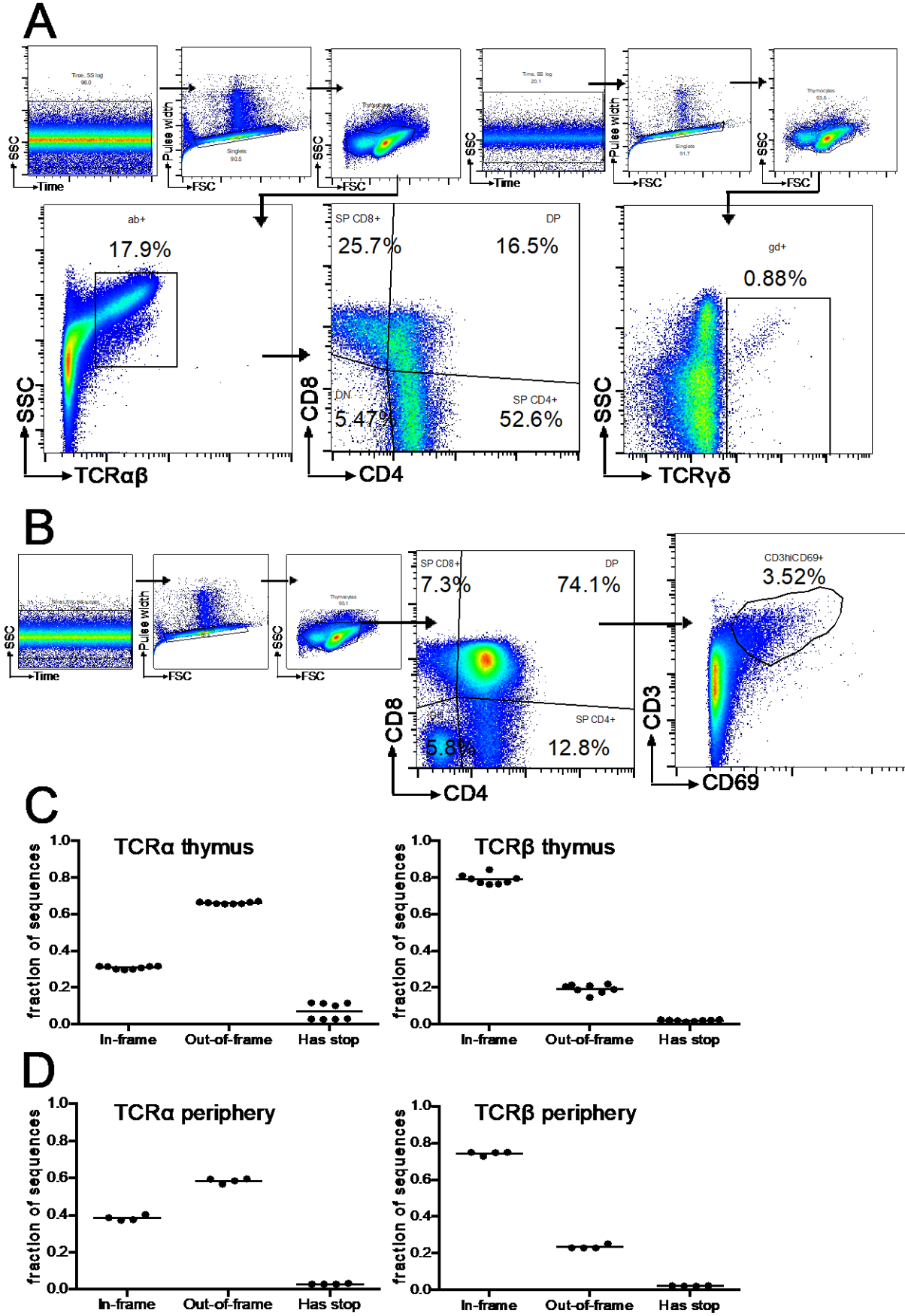
650 **Figure 6.** Sequence overlap among the high abundance clonotypes. Jaccard indexes (A) and non-
651 templated insertions in the shared in-frame sequences (B) among the top 1%, 2%, 5%, 10%, 20%
652 and 50% most abundant clonotypes and full repertoire in thymus samples A-D. The horizontal bars
653 show the average and error bars indicate the 95% confidence interval.

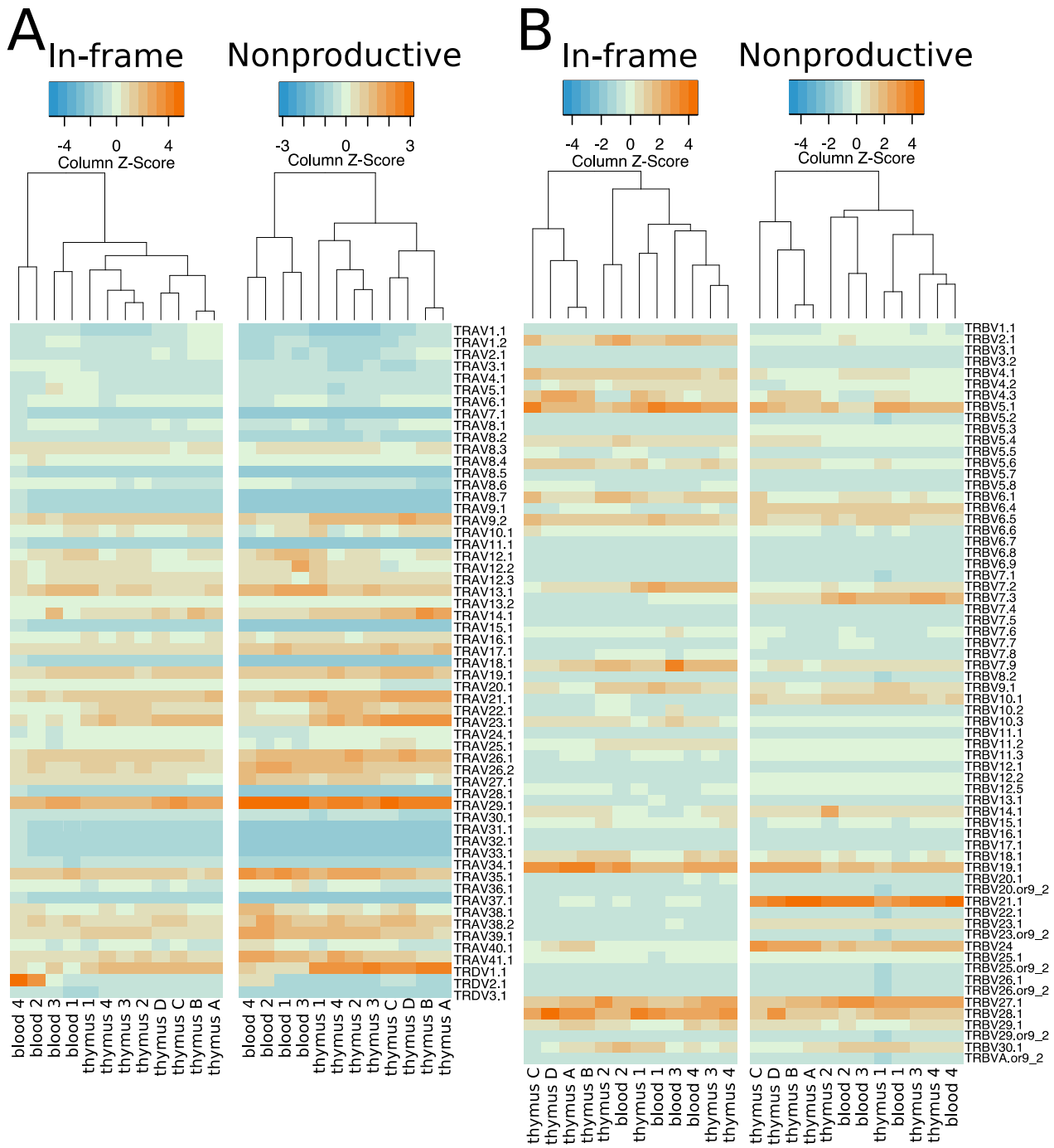
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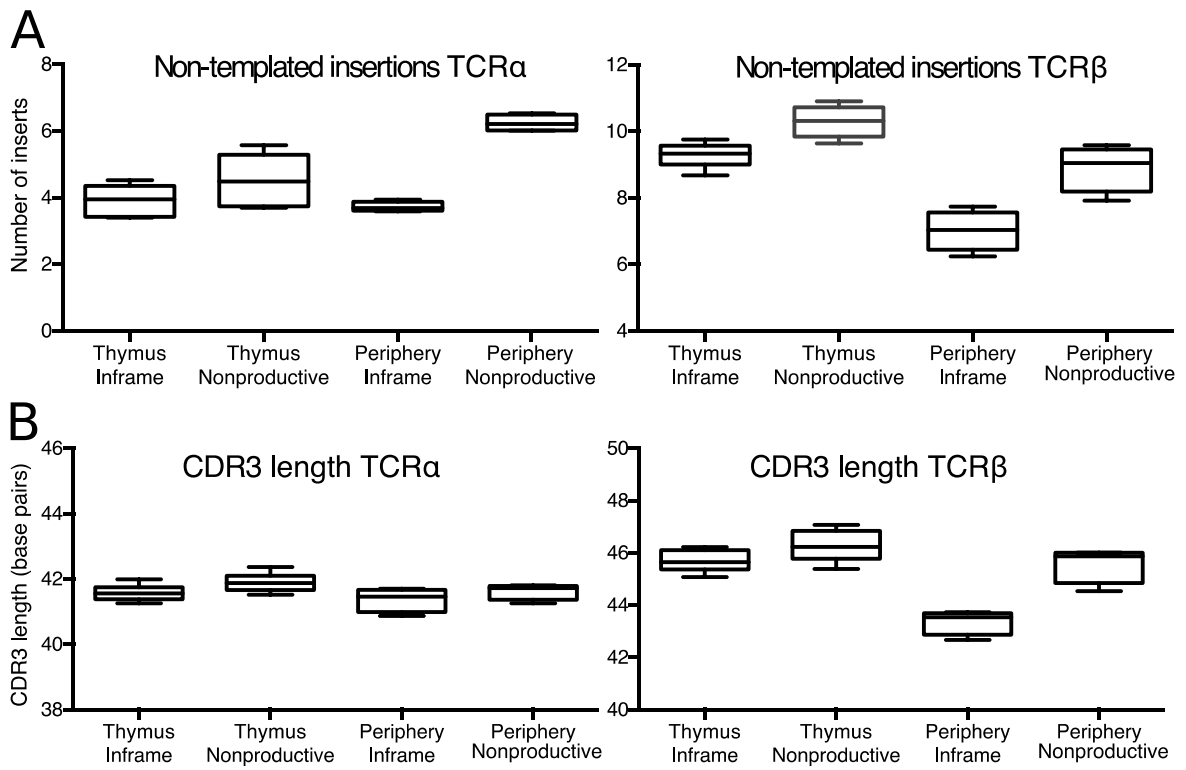
655 **Figure 7.** Sequence overlap between peripheral blood samples. Venn diagrams show the overlap of
656 unique in-frame clonotypes (A). Jaccard indexes (JI) for nonproductive, in-frame and amino acid
657 repertoires among unique clonotypes (B) and total genomes (C). The average JI and the 95%
658 confidence interval are displayed.

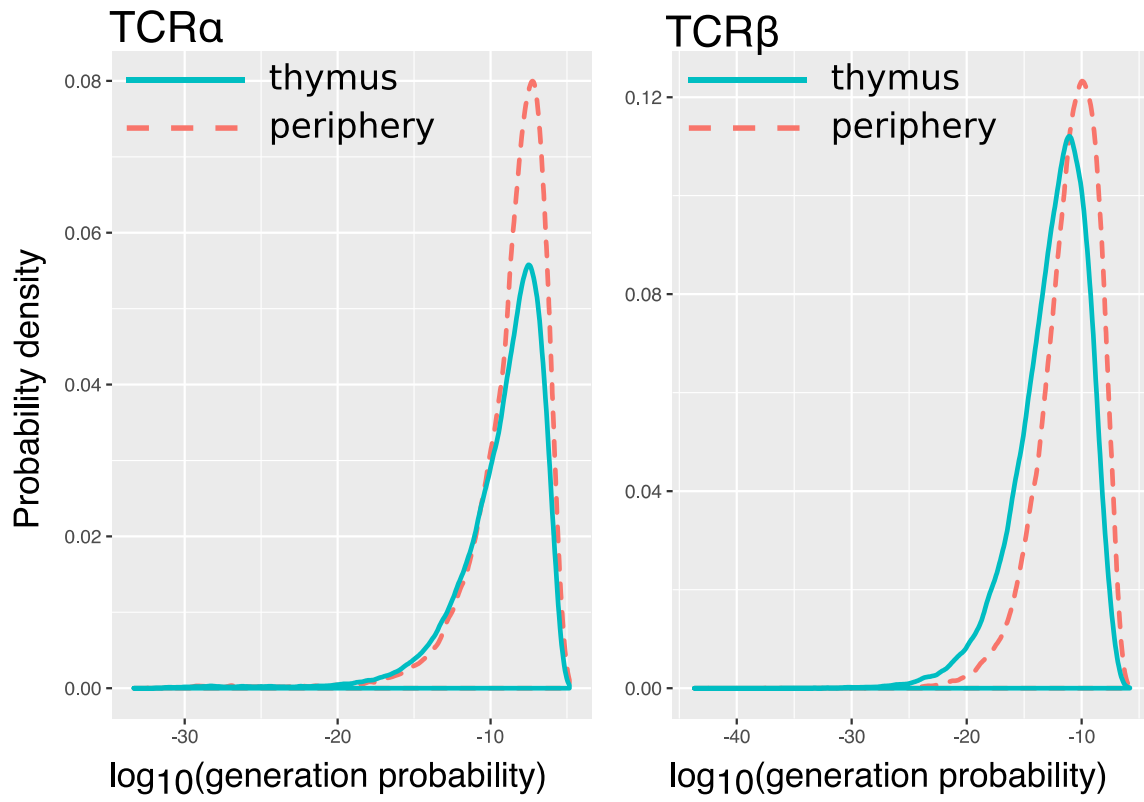
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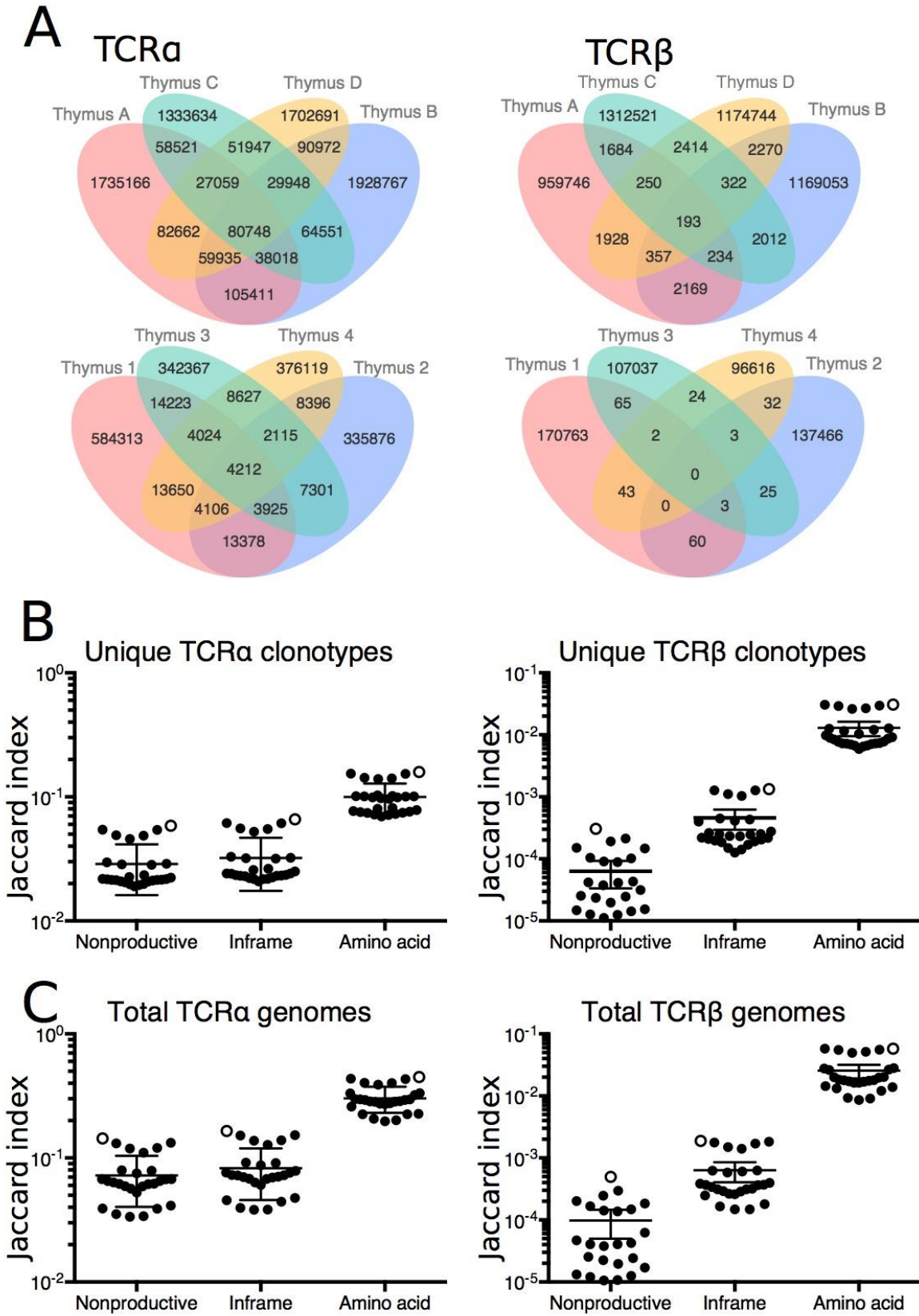
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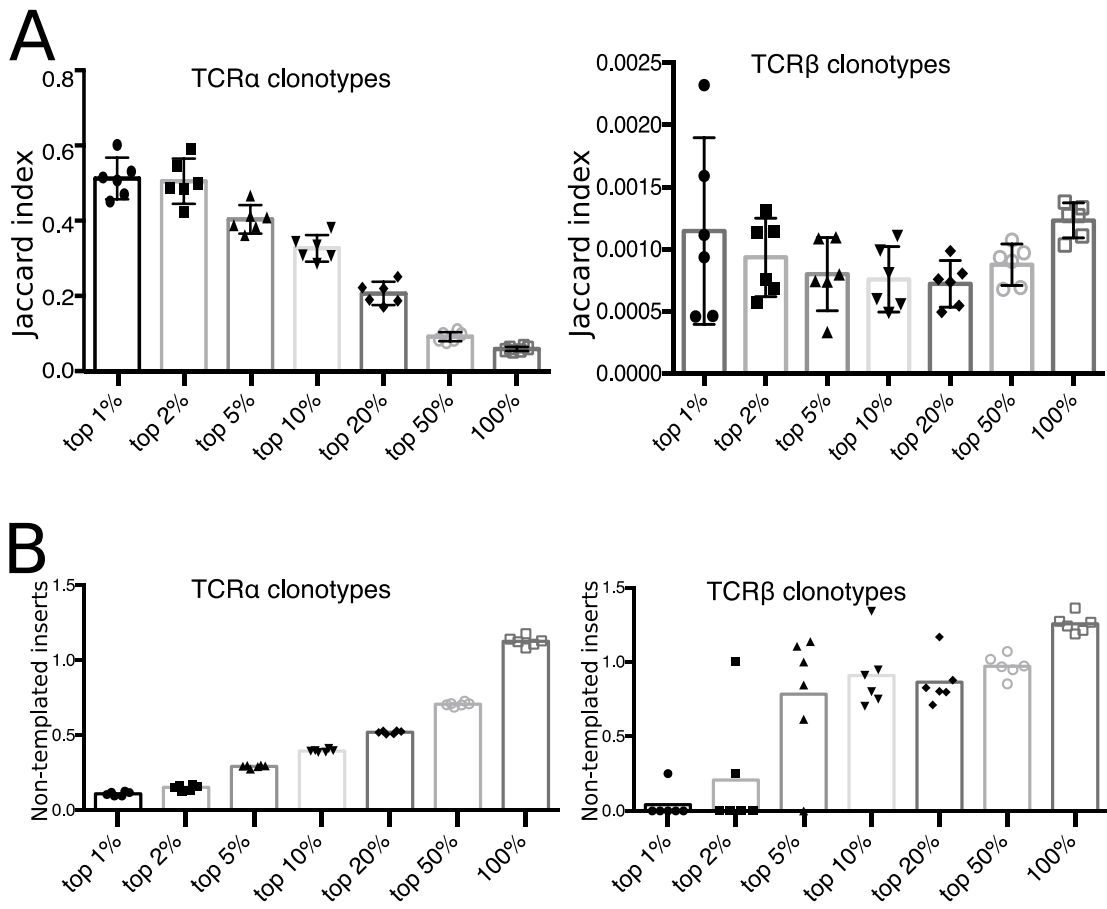




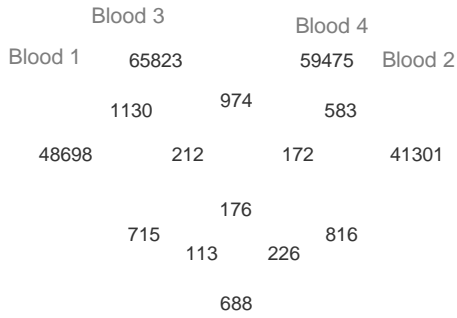








A TCR α



TCR β

